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August 11, 2005

BY FEDERAL EXPRESS

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

Attn: Dominique Delmas

Re: CHIRON CORPORATION

PCT Application No. PCT/US2005/006588 Our Ref.: 002441.00124:PP21431.004

Dear Sir:

Further to our letter dated August 1, 2005, we now enclose a certified copy of priority document for U.S. Serial No. 60/549,832 for the above-identified application.

Respectfully submitted,

Rebecca M. Hale

SCS/mh

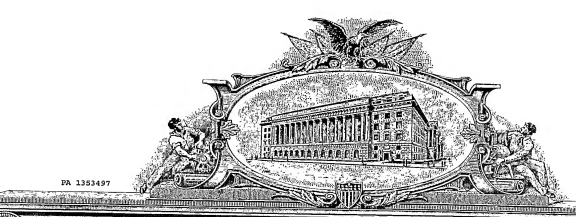
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August 09, 2005

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APPLICATION NUMBER: 60/549,832

FILING DATE: March 02, 2004

PA 1353497

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PTO/SB/16 (08-03)

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Docket Number 21431.001 INVENTOR(S)/APPLICANT(S) Residence Given Name (first and middle [if any]) Family or Surname (City and either State or Foreign Country) Giulio Ratti

Number 2 of 2

[Page 2 of 2]

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IMMUNOGENIC COMPOSITIONS FOR CHLAMYDIA PNEUNOMIAE All documents cited herein are incorporated by reference in their entirety.

Field

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5 The invention is in the field of immunology and vaccinology. In particular, it relates to immunogenic compositions comprising combinations of immunogenic molecules from *Chlamydia pneumoniae*.

Background Art

- The bacteria of the genus Chlamydia (and Chlamydophila, according to the recently 10 proposed but still controversial re-classification of Chlamydiaceae (Bush et al (2001) Int J Syst Evol Microbiol 51: 203-20; Everett et al (1999) Int J Syst Bacteriol 49: Pt2 415-40; Schachter et al (2001) Int J Syst Evol Microbiol 51: 249, 251-3) are obligate intracellular parasites of eukaryotic cells, which have a unique biphasic life cycle 15 involving two pleiomorphic developmental forms: an extracellular, metabolically inert, spore-like, infectious form (the elementary bodies, EBs) and an intracellular, noninfectious, replicative form (the reticulate bodies, RBs) which remains contained in a specialized cytoplasmic compartment (the Chlamydial inclusion). The EBs are responsible for the initial attachment to host cell surface and the establishment of the 20 cytoplasmic inclusion where EBs can differentiate to RBs and thus initiate the replicative stage. Eventually RBs revert to infectious EB forms able to start new replicative cycles in neighbouring host cells.
- As Chlamydia infection is an intracellular infection, the currently accepted paradigm is that effective anti-Chlamydial immunisation would require both an adequate T-cell response and high serum levels of neutralising antibodies and that "an ideal vaccine should induce long lasting (neutralising) antibodies and a cell mediated immunity that can quickly respond upon exposure to Chlamydia". Several sometimes contradictory studies have indicated that both CD4+ and CD 8 positive T cells have a role in Chlamydial clearance (Loomis and Starnback (2002) Curr Opin Microbiol 5: 87-91). Indeed, there now appears to be a prevailing consensus that specific CD4+ T cells and B cells are critical to the complete clearance of intracellular Chlamydia and for mediating recall immunity to Chlamydia infection (see Igietseme, Black and Caldwell (2002) Biodrugs 16: 19-35 and Igietseme et al (1999) Immunology 98: 510-519).

Whilst it is now possible to carry out searches of the whole Chlamydia pneumoniae genome, there is still insufficient information available on parallel proteome characterisation. By way of example, while sequence data is available for many of the Chlamydia pneumoniae antigens, there is insufficient characterisation of the Chlamydia antigens in terms of their immunological and/or biological function. By way of example, whilst applications such as WO 99/28475 and WO 99/27105 disclose sequence information, there is no characterisation of these sequences in terms of their immunological and/or biological function. In contrast, WO 02/02404 provides information on the immunogenicity and immunoaccessibility of certain Chlamydia proteins and highlights that (i) current genomic annotations and/or (ii) predictions based on cellular location and/or cellular function based on in-silico analyses may not always be accurate.

Applicants have recently engaged in a whole-genome search (Montigiani et al (2002) Infection and Immunity 70:368-379) for possible vaccine candidates among proteins potentially associated with the outer membrane of *C.pneumoniae*. For this study,

mouse antisera was prepared against over 100 recombinant His-tagged or Glutathione-S-transferase (GST) fusion proteins encoded by genes predicted by in silico analyses to be peripherally located in the *Chlamydial* cell. From this screening study, 53 recombinant proteins derived from the genome of *Chlamydial* (Chlamydophila pneumoniae (CPn) were described which induced mouse antibodies, capable of binding, in a FACS assay, to the surface of purified CPn cells.

The scope of the Montigiani study (*ibid*) was restricted to checking if polyclonal antisera produced in mice against the recombinantly expressed antibodies to CPn antigens were capable of binding to the surface of the CPn cells. No studies were carried out to test whether antisera against the recombinant FACS positive antigens were capable of interfering with EB *in vitro* infectivity of host cells – that is, whether the murine antibodies raised against the recombinantly expressed antigens could inhibit CPn infectivity *in vitro* to an extent greater than 50%, a property that common practice qualifies such antigens as "neutralising".

Indeed, so far, only few C. pneumoniae antigens with 'neutralizing' properties have been described in the literature: notably, a protein identified as 76-kDa-homolog protein (Perez-Melgosa et al (1994) Infect Immunity 62: 880-6), the surface-exposed outer membrane proteins MOMP (Wolf et al (2001) Infect Immun 69: 3082-91), PorB (Kawa et al (2002) J Immunol 168: 5184-91 and Kubo et al (2000) Mol Microbiol 38: 772-80), and very recently also the Pmp21 member of the Chlamydia-specific polymorphic family of outer membrane proteins (A.Szczepek, personal comunication). All these proteins were in fact selected in the earlier FACS-based screening study (Montigiani et al (2002) ibid). It can be however noted that outer membrane antigens, as it is the case for MOMP and PorB, could possibly present some kind of practical problems for a recombinant vaccine development project. For instance both MOMP and PorB are integral membrane proteins which appear to require a native conformation to maintain neutralizing epitopes which are discontinuous and conformation-dependent. The production of such proteins may require special processing steps (refolding) which could be undesirable in the preparation of an hypothetical vaccine. Other general problems may arise from the extent of allelic variation, and from regulated proteins which are not always expressed in all Chlamydial cell or all Chlamydial isolates.

Thus, it is desirable to provide improved compositions capable of eliciting an immune response upon exposure to *Chlamydia pneumoniae* proteins. It is also desirable to provide improved compositions comprising one or more combinations of two or more selected CPn proteins with complementary immunological and/or biological profiles capable of providing immunity against *Chlamydia*l induced disease and/or infection (such as in prophylactic vaccination) or (b) for the eradication of an established chronic *Chlamydia*l infection (such as in therapeutic vaccination).

Description of the Invention

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The present invention provides compositions comprising a first biological molecule from a *Chlamydia pneumoniae* bacterium and a second biological molecule from a *Chlamydia pneumoniae* bacterium. The term "biological molecule" includes proteins, antigens and nucleic acids. The compositions may also comprise further biological molecules preferably also from *Chlamydia pneumoniae*. That is to say, the compositions may comprise two or more biological molecules (eg. 3, 4, 5, 6, 7, 8 etc.)

at least two of which are from a *Chlamydia pneumoniae* bacterium (eg. 3, 4, 5, 6, 7, 8 etc.). Such compositions include those comprising (i) two or more different *Chlamydia pneumoniae* proteins; (ii) two or more different *Chlamydia pneumoniae* nucleic acids, or (iii) mixtures of one or more *Chlamydia pneumoniae* protein and one or more *Chlamydia pneumoniae* nucleic acid.

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In one preferred embodiment the first and second biological molecules are from different Chlamydia pneumoniae species (for example, from different Chlamydia pneumoniae serovars) but they may be from the same species. The biological molecules in the compositions may be from different serogroups or strains of the same 10 species. The first biological molecule is preferably selected from the group consisting of SEQ ID Nos 1-76. More preferably, it is selected from the group consisting of SEQ IDs 1-41 and/or SEQ ID Nos 42-76. It is preferably a purified or isolated biological molecule. The second biological molecule is preferably selected from the group consisting of SEQ ID Nos 1-76. More preferably, it is selected from the group · 15 consisting of SEQ IDs 1-41 and/or SEQ ID Nos 42-76. It is preferably a purified or isolated biological molecule. Specific compositions according to the invention therefore include those comprising: two or more biological molecules selected from the group consisting of SEQ ID Nos 1-41; one or more biological molecules selected from the group consisting of SEQ IDs 1-41 combined with one or more biological 20 molecules selected from the group consisting of SEQ IDs 42-76. One or both of the first and second biological molecules may be a Chlamydia pneumoniae biological molecule which is not specifically disclosed herein, and which may not have been identified, discovered or made available to the public or purified before this patent 25 application was filed.

A specific composition of the present invention may comprise a combination of *Chlamydia pneumoniae* antigens, said combination consisting of two, three, four, five or all six *Chlamydia pneumoniae* antigens of a first antigen group, said first antigen group consisting of: (1) pmp2; (2) pmp10; (3) Enolase; (4) OmpH-like protein; and (5) the products of CPn specific genes CPn0759 and CPn0042. These antigens are referred to herein as the 'first antigen group'.

Preferably, the composition of the invention comprises a combination of Chlamydia pneumoniae antigens, said combination selected from the group consisting of: (1) 35 pmp2 and pmp10; (2) pmp2 and Enolase; (3) pmp2 and OmpH-like protein; (4) pmp2 and CPn0759; (5) pmp2 and CPn0042; (6) pmp10 and Enolase; (7) pmp10 and OmpH-like protein; (8) pmp10 and CPn0759; (9) pmp10 and CPn0042; (10) Enolase and OmpH-like protein (11) Enolase and CPn0759; (12) Enolase and CPn0042; (13) OmpH-like protein and CPn0759 (14) OmpH-like protein and CPn0042; (15) 40 CPn0759 and CPn0042; (16) pmp2 and pmp10 and Enolase; (17) pmp2 and pmp10 and OmpH-like protein; (18) pmp2 and pmp10 and CPn0759; (19) pmp2 and pmp10 and CPn0042; (20) pmp2 and Enolase and OmpH-like protein; (21) pmp2 and Enolase and Cpn0759; (22) pmp2 and Enolase and CPn0042; (23) pmp2 and OmpHlike protein and CPn0759; (24) pmp2 and OmpH-like protein and CPn0042; (25) 45 pmp2 and Cpn0759 and CPn0042; and (26) pmp10 and Enolase and OmpH-like protein; (27) pmp10 and Enolase and CPn0759; (28) pmp10 and Enolase and CPn0042; (29) Enolase and OmpH-like protein and CPn0759; (30) Enolase and OmpH-like protein and CPn0042; (31) OmpH-like protein and CPn0759 and 50 CPn0042.

Preferably, the composition of *Chlamydia pneumoniae* antigens consists of pmp2, pmp10, Enolase, OmpH-like protein and CPn0759.

Preferably, the composition of *Chlamydia pneumoniae* antigens consists of pmp1, Enolase, OmpH-like protein and CPn0042.

Preferably, the composition of *Chlamydia pneumoniae* antigens consists of pmp2, pmp10, Enolase, OmpH-like protein and CPn0759 and CPn0042.

- The invention also provides for a slightly larger group of 12 Chlamydia pneumoniae antigens that are particularly suitable for immunisation purposes, particularly when used in combinations. (This second antigen group includes the six Chlamydia pneumoniae antigens of the first antigen group). These 12 Chlamydia pneumoniae antigens form a second antigen group of (1) pmp2; (2) pmp10; (3) Enolase; (4) OmpH-like protein; (5) CPn0759; (6) CPn0042; (7) ArtJ; (8) HtrA; (9) AtoS; (10) OmcA; (11) CPn0498; and (12) CPn0525. These antigens are referred to herein as the 'second antigen group'.
- The invention therefore provides a composition comprising a combination of Chlamydia pneumoniae antigens, said combination selected from the group consisting of two, three, four, five, six, seven, eight, nine, ten, eleven, or twelve Chlamydia pneumoniae antigens of the second antigen group. Preferably, the combination is selected from the group consisting of two, three, four or five Chlamydia pneumoniae antigens of the second antigen group. Still more preferably, the combination consists of six Chlamydia pneumoniae antigens of the second antigen group. Each of the Chlamydia pneumoniae antigens of the first and second antigen group are described in more detail below.

(1) Pmp10 (CPn0449)

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30 One example of a pmp10 protein is set forth as SEQ ID NO: 1 below (GenBank Accession No.GI:14195016). Preferred pmp10 proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 1; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 1, wherein n is 7 or more (e.g. 8, 10, 12, 14, 35 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These pmp2 proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 1. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 1. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino 40 acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 1. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). 45

SEO ID No 1

1 MKSQFSWLVL SSTLACFTSC STVFAATAEN IGPSDSFDGS TNTGTYTPKN TTTGIDYTLT
61 GDITLQNLGD SAALTKGCFS DTTESLSFAG KGYSLSFLNI KSSAEGAALS VTTDKNLSLT
121 GFSSLTFLAA PSSVITTPSG KGAVKCGGDL TFDNNGTILF KQDYCEENGG AISTKNLSLK
181 NSTGSISFEG NKSSATGKKG GAICATGTVD ITNNTAPTLF SNNIAEAAGG AINSTGNCTI

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241 TGNTSLVFSE NSVTATAGNG GALSGDADVT ISGNQSVTFS GNQAVANGGA IYAKKLTLAS
            301 GGGGGISFSN NIVQGTTAGN GGAISILAAG ECSLSAEAGD ITFNGNAIVA TTPQTTKRNS
           361 IDIGSTAKIT NLRAISGHSI FFYDPITANT AADSTDTLNL NKADAGNSTD YSGSIVFSGE
           421 KLSEDEAKVA DNLTSTLKQP VTLTAGNLVL KRGVTLDTKG FTQTAGSSVI MDAGTTLKAS
 5
           481 TEEVTLTGLS IPVDSLGEGK KVVIAASAAS KNVALSGPIL LLDNQGNAYE NHDLGKTQDF
           541 SFVQLSALGT ATTTDVPAVP TVATPTHYGY QGTWGMTWVD DTASTPKTKT ATLAWTNTGY
           601 LPNPERQGPL VPNSLWGSFS DIQAIQGVIE RSALTLCSDR GFWAAGVANF LDKDKKGEKR
           661 KYRHKSGGYA IGGAAQTCSE NLISFAFCQL FGSDKDFLVA KNHTDTYAGA FYIQHITECS
           721 GFIGCLLDKL PGSWSHKPLV LEGQLAYSHV SNDLKTKYTA YPEVKGSWGN NAFNMMLGAS
10
           781 SHSYPEYLHC FDTYAPYIKL NLTYIRQDSF SEKGTEGRSF DDSNLFNLSL PIGVKFEKFS
           841 DCNDFSYDLT LSYVPDLIRN DPKCTTALVI SGASWETYAN NLARQALQVR AGSHYAFSPM
           901 FEVLGQFVFE VRGSSRIYNV DLGGKFQF
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(2) Pmp2 = Polymorphic Outer Membrane Protein G Family (CPn 0013)

One example of a pmp2 protein is disclosed as SEQ ID NOs: 139 and 140 in WO 15 02/02606. {GenBank accession number: gi|4376270|gb|AAD18172.1 'CPn0013'; SEQ ID NO: 2 below}. Preferred pmp2 proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) 20 to SEQ ID NO: 2; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 1, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These pmp2 proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 2. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 1. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or 25 more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 2. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). 30

SEQ ID No 2

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	. 1	MKIPLRFLLI	SLVPTLSMSN	LLGAATTEEL	SASNSFDGTT	STTSFSSKTS
25	. 51	SATDGTNYVF	KDSVVIENVP	KTGETQSTSC	FKNDAAAGDL	NELGGGESET
35	101	FSNIDATTAS	GAAIGSEAAN	KTVTLSGFSA	LSFLKSPAST	WINGLOATHY
	151	KGNLSLLDND	KVLIQDNFST	GDGGATNCAG	STRIANMEST	CETCMCCCOMP
	201	GGAIHTKNLT	LSSGGETLFO	GNTADTAACK	CCATATADCC	TLSISGDSGD
	251	IIFEGNTIGA	TGTVSHSAID	T.CTCAVTMAT	DANCONTALADOG	TESTSGESGE
	301	CMADALMING	DUMODIMINA	TIGIDAKTIAL	RAAQGHTIYE	YDPITVTGST
40		DANDADIATIO	PDTGDNKEYT	GTIVFSGEKL	TEAEAKDEKN	RTSKLLQNVA
70	351	FKNGTVVLKG	DVVLSANGFS	QDANSKLIMD	LGTSLVANTE	SIELTNLEIN
	401	IDSLRNGKKI	KLSAATAQKD	IRIDRPVVLA	ISDESTYONG	FLNEDHSVDG
	451	ILELDAGKDI	VISADSRSID	AVOSPYGYOG	KWTTNWSTDD	KKYLACMYKO
	501	SFNPTAEOEA	PLVPNLLWGS	FIDURGEONE	TELCTECARY	KKAI VSWAKQ
	551	NVLHRSGREN	QRKFRHVSGG	VILLA CADMD	CODMICTORY	EVKLMAYGTZ
45	601	MNITATEARTYA	CCI DI OUDAG	AVVGASIRMP	GGDILSLGFA	QLFARDKDYF
•	651	THITTING IM	GSLRLQHDAS	PASAARTPPG	EGGLREILLP	YVSKTLPCSF
		YGQLSYGHTD	HRMKTESLPP	PPPTLSTDHT	SWGGYVWAGE	LGTRVAVENT
	701	SGRGFFQEYT	PFVKVQAVYA	RODSFVELGA	ISRDFSDSHI.	YNT ATPLCTE
	751	LEKRFAEQYY	HVVAMYSPDV	CRSNPKCTTT	LISNOGSWKT	KCCMI ADOAG
	801	IVOASGFRSL	GAAAELFGNF	CEEMBCCCDC	AND CONTR	MGOMINAKQAG
50				CACCOANGIO	INVUAGSKIK	F = .
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(3) Enolase (Cpn0800)

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One example of an 'Eno' protein is disclosed as SEQ ID NOs: 93 and 94 in WO 02/02606. {GenBank accession number: gi|4377111|gb|AAD18938.1| 'Cpn0800'; SEQ ID NO: 3 below}. Preferred Eno proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 2; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 2, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40,

50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These Eno proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 3. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 3. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 3. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

10 SEQ ID No 3

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15	101 151 201 251 301	GSPNKETLGA LINGGMHADN STGVGDEGGF YNVKTGTYDG LGEKVQIVGD	YQGKGVLQAV NAILGVSLAT GLEFQEFMIR APNLASNEEA RHYEEQIAIL DLFVTNPELI	KNVKEILFPL AHAAAATLRR PIGASSIKEA LELLLLAIEK SNLCDRYPID LEGISNGLAN	VKGCSVYEQS PLYRYLGGCF VNMGADVFHT AGFTPGKDIS SIEDGLAEED SVLIKENOIG	LKKLLHERGL LALDCAASSF YDGWALLTEV
20	3 3 T	LAQMAGYTTI RLMEIEEELG	ISHRSGETTD	TTTADLAVAG	NAGQIKTGSL	SRSERVAKYN

(4) OmpH-like outer membrane protein (CPn0301)

One example of 'OmpH-like' protein is disclosed as SEQ ID NOs: 77 & 78 in WO 02/02606. {GenBank accession number: gi|4376577|gb|AAD18450.1| 'CPn0301'; SEQ ID NO: 4 below}. Preferred OmpH-like proteins for use with the invention 25 comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 4; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 3, wherein n is 7 or more (e.g. 8, 10, 12, 14, 30 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmpH-like proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 4. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 4. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 19 or more, to remove the signal peptide) from the N-terminus of SEQ ID NO: 4. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 4

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MKKLLFSTFL LVLGSTSAAH ANLGYVNLKR CLEESDLGKK ETEELEAMKQ 1. QFVKNAEKIE EELTSIYNKL QDEDYMESLS DSASEELRKK FEDLSGEYNA YQSQYYQSIN QSNVKRIQKL IQEVKIAAES VRSKEKLEAI LNEEAVLAIA PGTDKTTEII AILNESFKKQ N* 151

(5) CPn0042 (Hypothetical)

One example of hypothetical protein is set forth as SEQ ID NO: 5 below. GenBank accession number: gi|4376296|gb|AAD18195.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 5; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 5, wherein n is 7 or

more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These Hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 5. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 5. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 19 or more, to remove the signal peptide) from the N-terminus of SEQ ID NO: 5 Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 5

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1 MEEVSEYLQQ	VENQLESCSK	RLTKMETFAL	GVRLEAKEET	ESTIT, SDUM	מדמות מחחדום
61 DMLSRVEEIE	RMLRMAELPL	LPIKEALTKA	FVOHNSCKEK	T.TYVGGUTYT	CDANTECEDIE
121 LQSLNQTLQR	AYKESOKVSG	LESEVENCER	OI VIOLIDORE	TILVEPIPE	SPAILTSEER
181 KESYHSERIH	VDCMDT.VEEV	VDDITOI BORD	ZUVDŽAKČI P	TOGVSLIKEE	LLFVTSTFRT
181 KFSYHSFRLH 241 ALRETEYWLY	DEEDNORMAN	IDDIDDERIK	ARWMAMSERY	RDAFQAFQEM	LKEGLVEEAQ

(6) CPn0795 (Hypothetical)

One example of hypothetical protein is disclosed as SEQ ID NOs: 63 & 64 in WO 02/02606. {GenBank accession number: gi|4377106|gb|AAD18933.1| 'CPn0795'; SEQ ID NO: 6 below). Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 25 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 6; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 6, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These 30 Hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 6. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 6. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 19 or more, to remove the signal peptide) from the N-terminus of SEQ ID NO: 6. Other 35 fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

40 SEQ ID No 6

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. 1	MKDLGTLGGT	SSTAKTVSPD	GKVIMGRSOT	ADGSWHAFMC	HTDFSSNNVL
51	FDLDNTYKTL	RENGRQLNSI	FNLONMMLOR	ASDHEFTERG	PSMIALCACI.
101	YVNALQNLPS	NLAAQYFGIA	YKIRPKYRLG	VFLDHNFSSH	VPNNENUSUM
151	RLWMGAFIGW	QDSDALGSSV	KVSFGYGKOK	ATITREOLEM	TEACCCPCUP
201	EGVAAQIEGR	YGKSLGGHVR	VOPFLGLOFV	HITRKEYTEN	αποπαπασυσ
251	TDYSTGVVYL	GIGSHIALVD	SLHVGTRMGM	EONFAAHTDR	FEGGTAGTON
301	FVFEKLDVTH	TRAFAEMRVN	YELPYLOSLN	LILRVNOOPL	OGVMGESSDI.
351	RYALGF*		_		2011101 00000

(7) ArtJ arginine periplasmic-binding protein (CPn 0482)

One example of 'ArtJ' protein is disclosed as SEQ ID NOs: 73 & 74 in WO 02/02606. {GenBank accession number: gi|4376767|gb|AAD18622.1| 'CPn0482'; SEQ ID NO: 7 below}. Preferred ArtJ proteins for use with the invention comprise an amino acid

sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 7; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 7, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These ArtJ proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 7. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 7. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 7. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). The ArtJ protein may be bound to a small molecule like arginine or another amino acid.

15 SEQ ID No 7

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1 MIKQIGRFFR AFIFIMPLSL TSCESKIDRN RIWIVGTNAT YPPFEYVDAQ
51 GEVVGFDIDL AKAISEKLGK QLEVREFAFD ALILNLKKHR IDAILAGMSI
101 TPSRQKEIAL LPYYGDEVQE LMVVSKRSLE TPVLPLTQYS SVAVQTGTFQ
151 EHYLLSQPGI CVRSFDSTLE VIMEVRYGKS PVAVLEPSVG RVVLKDFPNL
201 VATRLELPPE CWVLGCGLGV AKDRPEEIQT IQQAITDLKS EGVIQSLTKK
WQLSEVAYE*

(8) HtrA DO Serine Protease (CPn0979)

One example of an 'HrtA' protein is disclosed as SEQ ID NOs: 111 & 112 in WO 25 02/02606. {GenBank accession number: gi|4377306|gb|AAD19116.1| 'CPn0979'; SEQ ID NO: 8 below}. Preferred HrtA proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 8; and/or (b) which is a fragment of at least n consecutive amino acids 30 of SEQ ID NO: 8, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25; 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These HrtA proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 8. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 8. Other preferred 35 fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably at least 16 to remove the signal peptide) from the N-terminus of SEQ ID NO: 8. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic 40 domain, of a transmembrane domain, or of an extracellular domain). In relation to SEQ ID NO: 8, distinct domains are residues: 1-16; 17-497; 128-289; 290-381; 394-485; and 394-497.

SEQ ID No 8 45 MITKQLRSWL AVLVGSSLLA LPLSGQAVGK KESRVSELPQ DVLLKEISGG 51 FSKVATKATP AVVYIESFPK SQAVTHPSPG RRGPYENPFD YFNDEFFNRF 101 FGLPSQREKP QSKEAVRGTG FLVSPDGYIV TNNHVVEDTG KIHVTLHDGQ 151 KYPATVIGLD PKTDLAVIKI KSQNLPYLSF GNSDHLKVGD WAIAIGNPFG LQATVTVGVI SAKGRNQLHI ADFEDFIQTD AAINPGNSGG PLLNIDGQVI 201 50 GVNTAIVSGS GGYIGIGFAI PSLMANRIID QLIRDGQVTR GFLGVTLQPI 251 DAELAACYKL EKVYGALVTD VVKGSPADKA GLKQEDVIIA YNGKEVDSLS 301 351 MFRNAVSLMN PDTRIVLKVV REGKVIEIPV TVSQAPKEDG MSALQRVGIR 401 VQNLTPETAK KLGIAPETKG ILIISVEPGS VAASSGIAPG QLILAVNRQK VSSIEDLNRT LKDSNNENIL LMVSQGDVIR FIALKPEE* 451

(9) AtoS two-component regulatory system sensor histidine kinase protein (CPn0584)

One example of 'AtoS' protein is disclosed as SEQ ID NOs: 105 & 106 in WO 02/02606. GenBank accession number: gi|4376878|gb|AAD18723.1| 'CPn0584'; SEQ ID NO: 9 below}. Preferred AtoS proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 9; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 9, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 10 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These AtoS proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 9. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 9. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 9. Other fragments 15 omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 9

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- 1 MNVPDSKNLH PPAYELLEIK ARITQSYKEA SAILTATPDG ILLLSETGHF
 51 LICNSQAREI LGIDENLEIL NRSFTDVLPD TCLGFSIQEA LESLKVPKTL
 101 RLSLCKESKE KEVELFIRKN EISGYLFIQI RDRSDYKQLE NAIERYKNIA
 151 ELGKMTATLA HEIRNPLSGI VGFASILKKE ISSPRHQRML SSIISGTRSI
 201 NNLVSSMLEY TKSQPLNLKI INLQDFFSSL IPLLSVSFPN CKFVREGAQP
 251 LFRSIDPDRM NSVVWNLVKN AVETGNSPIT LTLHTSGDIS VTNPGTIPSE
 301 IMDKLFTPFF TTKREGNGLG LAEAQKIIRL HGGDIQLKTS DSAVSFFIII
- 30 (10) OmcA 9kDa-cysteine-rich lipoprotein(CPn0558)

diglyceride), and may thus have a N-terminal cysteine.

One example of 'OmcA' protein is disclosed as SEQ ID NOs: 9 & 10 in WO 02/02606. {GenBank accession number: gi|4376850|gb|AAD18698.1| 'CPn0558', 'OmcA', 'Omp3'; SEQ ID NO: 10 below}. Preferred OmcA proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 35 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 10; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 10, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmcA proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 10. Preferred fragments of (b) comprise an 40 epitope from SEQ ID NO: 10. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 18 or more to remove the signal peptide) from the N-terminus of SEQ ID NO: 10. Other fragments omit one or more domains of the protein (e.g. omission of a 45 signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). The protein may be lipidated (e.g. by a N-acyl

SEQ ID No 10

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1 MKKAVLIAAM FCGVVSLSSC CRIVDCCFED PCAPSSCNPC EVIRKKERSC 51 GGNACGSYVP SCSNPCGSTE CNSQSPQVKG CTSPDGRCKQ *

(11) CPn0498 (Hypothetical)

One example of a hypothetical protein is set forth as SEQ ID NO: 11 below. (GenBank Accession No. GI:4376784; AAD18638.1). Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 11; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 11, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These Hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 11. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 11. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 18 or more to remove the signal peptide) from the N-terminus of SEQ ID NO: 11. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). The protein may be lipidated (e.g. by a N-acyl diglyceride), and may thus have a N-terminal cysteine.

25 SEQ ID No 11

1 MNRRKARWVV ALFAMTALIS VGCCPWSQAK SRCSIDKYIP VVNRLLEVCG LPEAENVEDL 61 IESSSAWULT PEERFSGELV SICQVKDEHA FYNDLSLLHM TQAVPSYSAT YDCAVVFGGP 121 LPALRQRLDF LVREWQRGVR FKKIVFLCGE RGRYQSIEEQ EHFFDSRYNP FPTEENWESG 181 NRVTPSSEEE IAKFVWMQML LPRAWRDSTS GVRVTFLLAK PEENRVVANR KDTLLLFRSY 241 QEAFPGRVLF VSSQPFIGLD ACRVGQFFKG ESYDLAGPGF AQGVLKYHWA PRICLHTLAE

(12) CPn 0525 (hypothetical)

35 One example of 'Cpn0525' protein is disclosed as SEQ ID NOs: 117 & 118 in WO 02/02606. {GenBank accession number: gi|4376814|gb|AAD18665.1| 'CPn0525', SEQ ID NO: 12 below}. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 40 99.5% or more) to SEQ ID NO: 12; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 12, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmcA proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 12. Preferred fragments of (b) comprise an epitope from 45 SEQ ID NO: 12. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 18 or more to remove the signal peptide) from the N-terminus of SEQ ID NO: 12. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an 50 extracellular domain).

SEQ ID No 12

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- 1 MHDALLSILA IQELDIKMIR LMRVKKEHQK ELAKVQSLKS DIRRKVQEKE
 51 LEMENLKTQI RDGENRIQEI SEQINKLENQ QAAVKKMDEF NALTQEMTTA
 101 NKERRSLEHQ LSDLMDKQAG GEDLIVSLKE SLASTENSSS VIEKEIFESI
 151 KKINEEGKAL LEQRTELKHA TNPELLSIYE RLLNNKKDRV VVPIENRVCS
 201 GCHIVLTPQH ENLVRKKDRL IFCEHCSRIL YWQESQVNAQ ENSTAKRRRR
 251 RAAV*
- 10 Third Antigen Group

The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group. Such other *Chlamydia pneumoniae* antigens include a third antigen group consisting of (1) LcrE, (2) DnaK, (3) Omp85 homolog, (4) Mip-like; (5) OmcB (6) MurG (7) Cpn0186 and (8) fliY. These antigens are referred to herein as the "third antigen group".

(13) LcrE low calcium response E protein (CPn0324)

One example of a 'LcrE' protein is disclosed as SEQ ID NOs: 29 & 30 in WO 20 02/02606. {GenBank accession number: gi|4376602|gb|AAD18473.1| 'CPn0324'; SEQ ID NO: 13 below}. Preferred LcrE proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 13; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 13, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 25 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These LcrE proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 13. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 13. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 30 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 13. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

35 SEQ ID No 13

50

40 MAASGTGGL GGTQGVNLAA VEAAAAKADA AEVVASQEGS EMNMIQQG 51 LTNPAAATRT KKKEEKFQTL ESRKKGEAGK AEKKSESTEE KPDTDLAI ASGNSEISGQ ELRGLRDAIG DDASPEDILA LVQEKIKDPA LQSTALDY CONTROL OF TOP	OKY
40 101 ASGNSEISGQ ELRGLRDAIG DDASPEDILA LVQEKIKDPA LQSTALDY 151 QTTPPSQGKL KEALIQARNT HTEQFGRTAI GAKNILFASQ EYADQLNV 201 SGLRSLYLEV TGDTHTCDQL LSMLODRYTY ODMATUSSEL MYCMATHI	OKY
40 151 ASGNSEISGQ ELRGLRDAIG DDASPEDILA LVQEKIKDPA LQSTALDY 151 QTTPPSQGKL KEALIQARNT HTEQFGRTAI GAKNILFASQ EYADQLNV 201 SGLRSLYLEV TGDTHTCDQL LSMLODRYTY ODMAIUSSEL MYCMATHI	PT 77
201 SGLRSLYLEV TGDTHTCDQL LSMLODRYTY ODMAIUSSEL MYCMATRI	77.37
201 SGLRSLYLEV TGDTHTCDQL LSMLODRYTY ODMAIUSSEL MYCMATRI	
SGLKSLYLEV TGDTHTCDQL LSMLODRYTY ODMATUSET, MYCMARET	
251 OCCUPATION OF THE COLUMN ASSETS AND ASSETS OF THE COLUMN ASSETS OF T	/SP
251 OCDVIDGA OF CARRY	.VD
231 QUEIVESAUL OVIMTETENT, ONTERCONDE ECONOTES DO TEST	JACK.
251 QGPYVPSAQL QVLMTETRNL QAVLTSYDYF ESRVPILLDS LKAEGIQT	'PS
THE THEORY TASKVEREVE NUTGODINGS TO THE TERM OF THE TE	202
351 LROTSSRIES SADEPOOLCA MIANALDANA	SA
45 351 LRQTSSRLFS SADKRQQLGA MIANALDAVN INNEDYPKAS DFPKPYPW	IS*

(14) DnaK heat-shock protein 70 (chaperone) (CPn0503)

One example of 'DnaK' protein is disclosed as SEQ ID NO's: 103 & 104 in WO 02/02606. {GenBank accessionnumber: gi|4376790|gb|AAD18643.1| 'CPn0503'; SEQ ID NO: 14 below. Preferred DnaK proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 14; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 14, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30,

35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These DnaK proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 14. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 14. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 14. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

10 SEQ ID No 14

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					•	
	1	MSEHKKSSKI	IGIDLGTTNS	CVSVMEGGQA	KVITSSEGTR	TTPSTVAFKG
45	51	NEKLVGIPAK	RQAVTNPEKT	LGSTKRFIGR	KYSEVASEIQ	TVPYTVTSGS
	101	KGDAVFEVDG	KQYTPEEIGA	OILMKMKETA	EAYLGETVTE	AWTTWDAVEN
15	151	DSQRASTKDA	GRIAGLDVKR	IIPEPTAAAT.	AYGIDKVGDK	KINNEDICCO
	201	TFDISILEIG	DGVFEVLSTN	GDTLLGGDDF	DEVIIKWMIE	KINVEDIGGG
	251	SKDMMATODT	YDAABKAKER	TOGTOGER	DEVIINMILE	EFKKQEGIDL
		PICPAINTIQED	KDAMEKAKIE	LSGVSSTEIN	QPFITMDAQG	PKHLALTLTR
	301	AQFEKLAASL	IERTKSPCIK	ALSDAKLSAK	DIDDVLLVGG	MSRMPAVOET
00	351	VKELFGKEPN	KGVNPDEVVA	IGAAIOGGVL	GGEVKDVI.I.I.	DVTDLSLGTE
20	401	TLGGVMTTLV	ERNTTIPTOK	KOIFSTAADN	OPAUTTVITO	GEDDMY ADMA
	451	EIGRFDLTDI	PPAPRCHPOT	PUCEDIDANO	TEITIOSTES	COMPANDIA
	501	A COCK CERRE	a a superior of the superior o	PASEDIDMIG	TEHVSAKDVA	SGREQKIRIE
	201	ASSGLQEDEI	QRMVRDAEIN	KEEDKKRREA	SDAKNEADSM	IFRAEKATKD
	551	YKEQIPETLV	KEIEERIENV	RNALKDDAPT	EKTKEVTEDI.	SKAMOKICEG
	601	MQSQSASAAA	SSAANAKCCD	MITAUREUNT MARK	CECURADON	COCHRIGHTON
25	651	DIETTOR	~~~ MANGGE	MINITEDERKH	SESTKPPSNN	GSSEDHIEEA
	627	DVEIIDNDDK*				

(15) Omp85 homolog (Cpn0300)

One example of an Omp85 Homolog protein is disclosed as SEQ ID NOs: 147 & 148 in WO 02/02606. {GenBank accession number: gi|4376576|gb|AAD18449.1| 30 'CPn0300'; SEQ ID NO: 15 below}. Preferred Omp85 proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 15; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 15, wherein n is 7 or more (e.g. 8, 10, 35 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These DnaK proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 15. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 15. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or 40 one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 15. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 15

	1	MLIMRNKVIL	QISILALIQT	PLTLFSTEKV	KEGHVVVDSI	TITTEGENAS
EΛ	51	NKHPLPKLKT	RSGALFSQLD	FDEDLRILAK	EYDSVEPKVE	ESECKTNITAL.
50	101	HLIAKPSIRN	IHISGNQVVP	EHKILKTLQI	YRNDLFEREK	FLKGLDDLRT
	151	YYLKRGYFAS	SVDYSLEHNQ	EKGHIDVLIK	INEGPCGKIK	OLTESCISES
	201	EKSDIQEFIQ	TKQHSTTTSW	FTGAGLYHPD	IVEODSLAIT	NYLHNNGYAD
	251	AIVNSHYDLD	DKGNILLYMD	IDRGSRYTLG	HVHIOGFEVL	PKRLTEKOSO
EE	301	VGPNDLYCPD	KIWDGAHKIK	QTYAKYGYIN	TNVDVLFIPH	ATRETVOVTV
55	351	EVSEGSPYKV	GLIKITGNTH	TKSDVILHET	SLFPGDTFNR	LKLEDTEORI.
	401	RNTGYFQSVS	VYTVRSQLDP	MGNADQYRDI	FVEVKETTTG	NI-GLELGESS
	451	LDNLFGGIEL	SESNFDLFGA	RNIFSKGFRC	LRGGGEHLFL	KANEGDKVTD

	501	YTLKWTKPHF	LNTPWILGIE	LDKSINRALS	KDYAVOTYGG	MUSTTVTI.NE
	551	HLKYGLFYRG	SQTSLHEKRK	FLLGPNIDSN	KGFVSAAGVN	TAIVINGTIMEDD
	601	TPTTGIRGGV	TFEVSGLGGT	YHFTKLSLNS	SIYRKLTRKG	TLKTKGENOR
5	651	IKPYSNTTAE	GVPVSERFFL	GGETTVRGYK	SFIIGPKYSA	TEDOCCI.CCI.
3	701	LISEEFQYPL	IRQPNISAFV	FLDSGFVGLO	EYKISI.KDI.R	SSACECUPED
	751	VMNNVPVMLG	FGWPFRPTET	LNGEKIDVSQ	RFFFALGGMF	*

(16) Mip-like FKBP-type peptidyl-prolyl cis-trans (CPn0661)

10 One example of a Mip-like protein is disclosed as SEQ ID NOs: 55 & 56 in WO 02/02606. {GenBank accession number: gi|4376960|gb|AAD18800.1| 'CPn0661'; SEQ ID NO: 16 below}. Preferred Mip-like proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 16; and/or (b) which is a fragment of at least n15 consecutive amino acids of SEQ ID NO: 16, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These miplike proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 16. Preferred fragments of (b) comprise an epitope from 20 SEQ ID NO: 16. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 16. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an 25 extracellular domain).

SEQ ID No 16

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30	1	MNRRWNLVLA	TVALALSVAS	CDVRSKDKDK	DOGSLVEYKD	NKDTNDTELS
	51	DNQKLSRTFG	HLLARQLRKS	EDMFFDIAEV	AKGLOAELVC	KSAPI.TETEV
	101	EEKMAEVQKL	VFEKKSKENL	SLAEKFLKEN	SKNAGVVEVO	PSKT-OVKTTV
	151	EGAGKAISGK	PSALLHYKGS	FINGOVESSS	EGNNEPTIJP	LCOTT DCFAT.
25	201	GMQGMKEGET	RVLYIHPDLA	YGTAGOLPPN	SLLIFEINLT	OASADEWAAV
35	251	PQEGNQGE*				X. 101 TO TO ALVE A

(17) OmcB 60 kDa Cysteine rich OMP (CPn0557)

One example of an OmcB protein is disclosed as SEQ ID NOs: 47 & 48 in WO 02/02606. {GenBank accession number: gi|4376849|gb|AAD18697.1| 'CPn0557'; SEQ ID NO: 17 below}. Preferred OmcB proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 17; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 17, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmcB proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 17. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 17. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 17. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain)..

SEQ ID No 17

		1	MSKLIRRVVT	VLALTSMASC	FASGGIEAAV	AESLITKIVA	SAETKPAPVP
_		51	MTAKKVRLVR	RNKQPVEQKS	RGAFCDKEFY	PCEEGRCOPV	ENOUESCACE
5	•	101	TARAKANDDC	NVEICQSVPE	YATVGSPYPI	EILAIGKKDC	WOWT TOOT. D
		151	CEAEFVSSDP	ETTPTSDGKL	VWKIDRLGAG	DKCKTTVWVK	DI.KEGCGETA
•		201	ATVCACPELR	SYTKCGOPAI	CIKQEGPDCA	CLRCPVCVKT	ENTANTECENTA
		251	RNVTVDNPVP	DGYSHASGOR	VLSFNLGDMR	PGDKKVETVE	ECDODDCOIM
4.0		301	NVATVTYCGG	HKCSANVTTV	VNEPCVQVNI	SCADWSYVCK.	PURVETCURA
10		351	PGDLVLHDVV	IODTLPSGVT	VLEAPGGEIC	CMKAVMBIRE	MCDCEMI OHA
		401	LVVKAQVPGR	FTNOVAVTSE	SNCGTCTSCA	ETTTHWKCIA	MCEGETTÖLK
		451	DPICVGENTV	YRICVTNRGS	AEDTNVSLIL	KECKET ODIN	WINDCARDIN
		501	GNTVVFDALP	KLGSKESVEF	SVTLKGIAPG	DADGEATICS	DOLUCIA
		551	ENTHVY*		~ · · · · · · · · · · · · · · · · · · ·	DUMGENTIDD	DITTPLASDL
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(18) MurG peptidoglycan transferase protein (CPn0904)

One example of a 'MurG' protein is disclosed as SEQ ID NOs: 107 & 108 in WO 02/02606. {GenBank accession number: gi|4377224|gb|AAD19042.1| 'CPn0904'; SEQ ID NO: 18 below}. Preferred MurG proteins for use with the invention 20 comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 18; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 18, wherein n is 7 or more (e.g. 8, 10, 12, 14, 25 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These MurG proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 18. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 18. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino 30 acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 18. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). The MurG may be lipidated e.g. with undecaprenyl.

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SEQ ID No 18

40	. 101 151 201	QGISYREIPS GSYHSLPVLL KHFRCPAEEV PQALVKLVNK	GLPTVLNPIK AGLSHKIPLF FLPKRSFSLG YPNMYVHHIV	IMSRTLSLCS	GYLKARKELK VNQLFSRYAR TPTICVVGGS HVYNRGEVLC	GIGVNFSPVT QGAQILNTCV CVKPFFFOLL
45	301	LEGGTMILEK AFICECL*	ELTEKLLVEK	VTFALDSHNR	EKQRNSLAAY	SQQRSTKTFH

(19) CPn0186 (Hypothetical)

One example of a hypothetical protein is set forth as SEQ ID NO: 19 below}. (GenBank Accession No. GI:4376456; AAD18339.1). Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 19; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 19, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants,

homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 19. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 19. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 19. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 19

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ı	v

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1	MSSPVNNTPS	APNIPIPAPT	TPGIPTTKPR	SSETEKVITY	AKVII.ENINA	TSGALGTILG
61	LSGALTPGTG	TALLUTERUC	MITT TOT TE VO	OTOGGGGGGG	MILTIDESTAN	NORLTVITTT
101	7.500.000.000	THEOTER	MATITICATIVA	SISGGEERKL	REEVSRFTSE	NQRLTVITTT
727	LETEVKDLKA	AKDQLTLEIE	AFRNENGNLK	TTAEDLEEOV	SKLSEOLEAL	ERINQLIQAN
181	AGDAOEISSE	LKKLISGWDS	KWESTNEST	ONTENTACOR	MIOUS OFFI	AMQEQIQALQ
241	A PETT CAMPAGO	mar over	TOTAL STATES	SWITKATTPGÖF	MAGEWGIHAK	AMQEQIQALQ
241	VETTIGMUNDS	TALQKSVENL	LVQDQALTRV	VGELLESENK	LSOACSALRO	EIEKLAQHET
301	SLQQRIDAML	ACEONLAEOV	TALEKMKOEA	OKAESERTAC	VEDETECTE	TPPPTTPVVE
361	GDESOFEDEG	GTDDVGCDGG	PVDRATCDCO	K-emont tue	AMDITTGREE	TEEFTIEVAE

(20) FliY Glutamine Binding Protein (CPn0604)

One example of a hypothetical protein is set forth as SEQ ID NOs: 11 & 12 in WO 20 02/02606. {GenBank accession number: gi|4376900|gb|AAD18743.1| 'CPn0604'; SEQ ID NO: 20 below). Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 20; and/or (b) which is a fragment of at least n25 consecutive amino acids of SEQ ID NO: 20, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 20. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 20. Other preferred fragments lack one or more amino 30 acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 20. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a 35 transmembrane domain, or of an extracellular domain).

SEQ ID No 20

40	2.7	DINAFFNDFA	SEINYKENLN	INIVNODWVH	LFENLDDKKT	FPKQFGIYTS QGAFTSVLPT VYKFDSSVLV
	121	AQNIPDAVIS	LYQHVPIALE	ALTSNCYDAL	LAPVIEVTAL	IETAYKGRLK DATKOPVDI.D

The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group or the third antigen group. Such other *Chlamydia pneumoniae* antigens include a fourth antigen group consisting one or more members of the PMP family. These antigens are referred to herein as the "fourth antigen group". Each of the *Chlamydia pneumoniae* antigens of the fourth antigen group is described in more detail below.

Fourth Antigen Group

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(21) Polymorphic Membrane Proteins (PMP)

A family of twenty one Chlamydia pneumoniae genes encoding predicted polymorphic membrane proteins (PMP) have been identified (pmp1 to pmp21).

15 Pmp1 (CPn0005)

One example of a Pmp1 protein is set forth as SEQ ID NOs: 41 & 42 in WO 02/02606. {GenBank accession number: gi|4376260|gb|AAD18163.1 'CPn0005'; SEQ ID NO: 21 below}.

20 SEQ ID No 21

	_					
	1	MRFSLCGFPL	VFSFTLLSVF	DISLIBATTIS	LTPEDSFHGD	SONAERSYNV
	51	QAGDVYSLTG	DVSISNVDNS	ALNKACFNVT	SGSVTFAGNH	HGLYFNNISS
05	101	GTTKEGAVLC	CQDPQATARF	SGFSTLSFIO	SPGDIKEOGC	LYSKNALMI.I.
25	151	NNYVVRFEQN	QSKTKGGAIS	GANVTIVGNY	DSVSFYQNAA	TECCATHEEC
	201	PLQIAVNQAE	IRFAONTAKN	GSGGALYSDG	DIDIDQNAYV	I.PDPMPAI.TO
	251	AIGKGGAVCC	LPTSGSSTPV	PIVTESDNKO	LVFERNHSIM	CCCATVADVI
	301	SISSGGPTLF	TNNTSYANSO	NT.CCATATOT	GGEISLSAEK	GGGATIAKKL
	351	SLPFLNGTHI	I ONAKET KTO	ADMOVETERY	DPITSEADGS	GITTFQGNRT
30 ·	401	NKEYTGTTLE	SCERCI AND	WWG121FL1	DPITSEADGS	TQLNINGDPK
	451	FTOCDCCUTY	T DT COURT TAC	RDFRSTIPQN	VNLSAGYLVI	KEGAEVTVSK
	501	TIQSEGSILLV	DDLGIKLIAS	KEDIAITGLA	IDIDSLSSSS	TAAVIKANTA
		NKQISVIDSI	ELISPTGNAY	EDLRMRNSQT	FPLLSLEPGA	GGSVTVTAGD
	551	FLPVSPHYGF	QGNWKLAWTG	TGNKVGEFFW	DKINYKPRPE	KECMI VIDNITI.
0.5	601	WGNAVDVRSL	MQVQETHASS	LOTDRGLWID	GIGNFFHVSA	SEDNIRVEHM
35	651	SGGYVLSVNN	EITPKHYTSM	AFSOLFSRDK	DYAVSNNEYR	MVI.GGVI.VOV
	701	TTSLGNIFRY	ASRNPNVNVG	ILSERFLOND	LMIFHFLCAY	CHYMANIAMA
	751	YANFPMVKNS	WRNNOWATEC	CCCMDITATE	NGRLFQGAIP	GUAINDMKID
	801	OCDEKETTAD	CDDECMOOLE	GEOMETIC SEE	MGKLFQGAIP	FMKTÖTATÄ
	851	TEDADDOGEN	GRAFANGSLI	STSARTGIKE	EKLALSQDVL	YDFSFSYIPD
40		IFKADPSCEA	ALVISGDSWL	VPAAHVSRHA	FVGSGTGRYH	FNDYTELLCR
70	901	GSIECRPHAR	NYNINCGSKF	RF*	•	

Pmp 4 (CPn0017)

One example of a Pmp 4 protein is designated SEQ ID NO: 22. The sequence for pmp4 protein can be found at AE001587.1 GI:4376271.

Pmp 6 (CPn0444)

One example of a Pmp 6 protein is set forth as SEQ ID NO^s 31 & 32 in WO 02/02606. {GenBank accession number: gi|4376727|gb|AAD18588.1| 'CPn0444'; SEQ ID NO: 23 below}.

SEQ ID No 23

EE	1	MKYSLPWLLT	SSALVFSLHP	LMAANTDLSS	SDNYENGSSG	SAAFTAKETS
55	51	DASGTTYTLT	SDVSITNVSA	ITPADKSCFT	NTGGALSEVG	ADHST MACOUT
	101	ALTHDGAAIN	NTNTALSESG	FSSLLTDSAP	ATGTSGGKGA	TOTOTIVECOM

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151 ATFTDNASVT LQKNTSEKDG AAVSAYSIDL AKTTTAALLD QNTSTKNGGA
                     LCSTANTTVQ GNSGTVTFSS NTATDKGGGI YSKEKDSTLD ANTGVVTFKS
                251
                     NTAKTGGAWS SDDNLALTGN TQVLFQENKT TGSAAQAMNP EGCGGAICCY
                     LATATOKTGL AISQNQEMSF TSNTTTANGG AIYATKCTLD GNTTLTFDQN
 5
                351
                     TATAGCGGAI YTETEDFSLK GSTGTVTFST NTAKTGGALY SKGNSSLTGN
                     TNLLFSGNKA TGPSNSSANQ EGCGGAILAF IDSGSVSDKT GLSIANNQEV
                401
                451
                     SLTSNAATVS GGAIYATKCT LTGNGSLTFD GNTAGTSGGA IYTETEDFTL
                     TGSTGTVTFS TNTAKTGGAL YSKGNNSLSG NTNLLFSGNK ATGPSNSSAN
                501
                     QEGCGGAILS FLESASVSTK KGLWIEDNEN VSLSGNTATV SGGAIYATKC
                551
10
                601
                    ALHGNTTLTF DGNTAETAGG AIYTETEDFT LTGSTGTVTF STNTAKTAGA
                     LHTKGNTSFT KNKALVFSGN SATATATTTT DQEGCGGAIL CNISESDIAT
                651
                701
                     KSLTLTENES LSFINNTAKR SGGGIYAPKC VISGSESINF DGNTAETSGG
                    AIYSKNLSIT ANGPVSFTNN SGGKGGAIYI ADSGELSLEA IDGDITFSGN
                751
                    RATEGTSTPN SIHLGAGAKI TKLAAAPGHT IYFYDPITME APASGGTIEE
                801
15
                851
                    LVINPVVKAI VPPPQPKNGP IASVPVVPVA PANPNTGTIV FSSGKLPSQD
                    ASIPANTTTI LNQKINLAGG NVVLKEGATL QVYSFTQQPD STVFMDAGTT
                901
                951
                    LETTTTNNTD GSIDLKNLSV NLDALDGKRM ITIAVNSTSG GLKISGDLKF
                    HNNEGSFYDN PGLKANLNLP FLDLSSTSGT VNLDDFNPIP SSMAAPDYGY
               1001
                     QGSWTLVPKV GAGGKVTLVA EWQALGYTPK PELRATLVPN SLWNAYVNIH
               1051
20
               1101
                    SIQQEIATAM SDAPSHPGIW IGGIGNAFHQ DKQKENAGFR LISRGYIVGG
                    SMTTPQEYTF · AVAFSQLFGK SKDYVVSDIK SQVYAGSLCA QSSYVIPLHS
               1151
               1201
                    SLRRHVLSKV LPELPGETPL VLHGQVSYGR NHHNMTTKLA NNTQGKSDWD
               1251
                    SHSFAVEVGG SLPVDLNYRY LTSYSPYVKL QVVSVNQKGF QEVAADPRIF
                    DASHLVNVSI PMGLTFKHES AKPPSALLLT LGYAVDAYRD HPHCLTSLTN
               1301
25
              1351
                    GTSWSTFATN LSRQAFFAEA SGHLKLLHGL DCFASGSCEL RSSSRSYNAN
              1401
                    CGTRYSF*
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Pmp 7 (CPn0445)

One example of a Pmp 7 protein is set forth as SEQ ID NOs 153 & 154 in WO 02/02606. {GenBank accession number: gi|4376728|gb|AAD18589.1| 'CPn0445'; SEQ ID NO: 24 below}.

SEQ ID No 24

35

	•				
1	MKSSVSWLFF	SSIPLFSSLS	IVAAEVTLDS	SNNSYDGSNG	TTFTVFSTTD
51	AAAGTTYSLL	SDVSFQNAGA	LGIPLASGCF	LEAGGDLTFO	GNOHALKEAE
101	INAGSSAGTV	ASTSAADKNL	LFNDFSRLSI	ISCPSLLLSP	TGOCALKSVG
151	NLSLTGNSQI	IFTONFSSDN	GGVINTKNFL	LSGTSOFASE	SRNOÄETGKO
201	GGVVYATGTI	TIENSPGIVS	FSONLAKGSG	GALYSTONCS	TTDNFOVTED
251	GNSAWEAAQA	QGGAICCTTT	DKTVTLTGNK	NLSFTNNTAL	TYGGATSGLK
301	VSISAGGPTL	FOSNISGSSA	GOGGGGAINI	ASAGELALSA	TSGDTTENMN
351	QVTNGSTSTR	NAINIIDTAK	VTSIRAATGO	SIYFYDPITN	PGTAASTOTT.
401	NLNLADANSE	IEYGGAIVFS	GEKLSPTEKA	IAANVTSTTR	OPAVI.ARCDI.
451	VLRDGVTVTF	KDLTOSPGSR	ILMDGGTTLS	AKEANLSING	LAVNILSSING
501	TNKAALKTEA	ADKNISLSGT	IALIDTEGSF	YENHNLKSAS	TYPLLELTTA
551	GANGTITLGA	LSTLTLQEPE	THYGYOGNWO	LSWANATSSK	IGSINWTRTG
601	YIPSPERKSN	LPLNSLWGNF	IDIRSINOLI	ETKSSGEPFE	RELWISCIAN
651	FFYRDSMPTR	HGFRHISGGY	ALGITATTPA	EDOLTFAFCO	LEARDRNHIT
701	GKNHGDTYGA	SLYFHHTEGL	FDIANFLWGK	ATRAPWVISE	TSOTTDISED
751					
801	EVEPFVKVOY	IYAHOODFYE	RHAEGRAFNK	SELTIMETET	CVTPPPPDCVC
851	EKGTYDLTLM	YILDAYRRNP	KCOTSLIASD	ANWMAYCTMI.	ADUCECTOVY
901	NHFOVNPHME	IFGOFAFEVR	SSSRNYNTNT	GSKECE*	THE SAKEM
				JU112 CX	
	51 101 151 201 251 301 351 401 451 501 551 601 651 701 751 801 851	51 AAAGTTYSLL 101 INAGSSAGTV 151 NLSLTGNSQTV 201 GGVVYATGTI 251 GNSAWEAAQA 301 VSISAGGPTL 351 QVTNGSTSTR 401 NLNLADANSE 451 VLRDGVTVTF 501 TNKAALKTEA 551 GANGTITLGA 601 YIPSPERKSN 651 FFYRDSMPTR 701 GKNHGDTYGA 751 AKFSYLHTDN 801 EVEPFVKVQY 851 EKGTYDLTLM	51 AAAGTTYSLL SDVSFQNAGA 101 INAGSSAGTV ASTSAADKNL 151 NLSLTGNSQI IFTQNFSSIN 201 GGVVYATGTI TIENSPGIVS 251 GNSAWEAAQA QGGAICCTTT 301 VSISAGGPTL FQSNISGSSA 351 QVTNGSTSTR NAINIIDTAK 401 NLNLADANSE IEYGGAIVFS 451 VLRDGVTVTF KDLTQSPGSR 501 TNKAALKTEA ADKNISLSGT 551 GANGTITLGA LSTLTLQEPE 601 YIPSPERKSN LPLNSLWGNF 651 FFYRDSMPTR HGFRHISGGY 701 GKNHGDTYGA SLYFHHTEGL 751 AKFSYLHTDN HMKTYYTDNS 801 EVEPFVKVQY IYAHQQDFYE 851 EKGTYDLTLM YILDAYRNP	51 AAAGTTYSLL SDVSFQNAGA LGIPLASGCF 101 INAGSSAGTV ASTSAADKNL LFNDFSRLSI 151 NLSLTGNSQI IFTQNFSSDN GGVINTKNFL 201 GGVVYATGTI TIENSPGIVS FSQNLAKGSG 251 GNSAWEAAQA QGGAICCTTT DKTVTLTGNK 301 VSISAGGPTL FQSNISGSSA GQGGGGAINI 351 QVTNGSTSTR NAINIIDTAK VTSIRAATGQ 401 NLNLADANSE IEYGGAIVFS GEKLSPTEKA 451 VLRDGVTVTF KDLTQSPGSR ILMDGGTTLS 501 TNKAALKTEA ADKNISLSGT IALIDTEGSF 551 GANGTITLGA LSTLTLQEPE THYGYQGNWQ 601 YIPSPERKSN LPLNSLWGNF IDIRSINQLI 651 FFYRDSMPTR HGFRHISGGY ALGITATTPA 701 GKNHGDTYGA SLYFHHTEGL FDIANFLWGK 751 AKFSYLHTDN HMKTYYTDNS IIKGSWRNDA 801 EVEPFVKVQY IYAHQQDFYE RHAEGRAFNK 851 EKGTYDLTLM YILDAYRRNP KCQTSLIASD	51 AAAGTTYSLL SDVSFQNAGA LGIPLASGCF LEAGGDLTFQ 101 INAGSSAGTV ASTSAADKNL LFNDFSRLSI ISCPSLLLSP 151 NLSLTGNSQI IFTQNFSSDN GGVINTKNFL LSGTSQFASF 201 GGVYYATGTI TIENSPGIVS FSQNLAKGSG GALYSTDNCS 251 GNSAWEAAQA QGGAICCTTT DKTVTLITGNK NLSFTNNTAL 301 VSISAGGPTL FQSNISGSSA GQGGGGAINI ASAGELALSA 351 QVTNGSTSTR NAINIIDTAK VTSIRAATGQ SIYFYDPITN 401 NLNLADANSE IEYGGAIVFS GEKLSPTEKA IAANVTSTIR 451 VLRDGVTVTF KDLTQSPGSR ILMDGGTTLS AKEANLSING 501 TNKAALKTEA ADKNISLSGT IALIDTEGSF YENHNLKSAS 551 GANGTITLGA LSTLTLQEPE THYGYQGNWQ LSWANATSSK 601 YIPSPERKSN LPLNSLWGNF IDIRSINQLI ETKSSGEPFE 651 FFYRDSMPTR HGFRHISGGY ALGITATTPA EDQLTFAFCQ 651 FFYRDSMPTR HGFRHISGGY ALGITATTPA EDQLTFAFCQ 701 GKNHGDTYGA SLYFHHTEGL FDIANFLWGK ATRAPWVLSE 751 AKFSYLHTDN HMKTYYTDNS IIKGSWRNDA FCADLGASLP 801 EVEPFVKVQY IYAHQQDFYE RHAEGRAFNK SELINVEIPI 851 EKGTYDLTLM YILDAYRNP KCQTSLIASD ANWMAYGTNL

Pmp 8 (CPn0446)

One example of a Pmp 8 protein is set forth as SEQ ID NO^s 45 & 46 in WO 02/02606. {GenBank accession number: gi|4376729|gb|AAD18590.1| 'CPn0446'; SEQ ID NO: 25 below}.

SEQ ID No 25

	1	MKIPLHKLLI	SSTLVTPILL	SIATYGADAS	LSPTDSFDGA	GGSTFTPKST
5	51	ADANGTNYVL	SGNVYINDAG	KGTALTGCCF	TETTGDLTFT	CKCVCFCFNTP
5	101	VDAGSNAGAA	ASTTADKALT	FTGFSNLSFI	AAPGTTVASG	KSTT. CCACAT.
	151	NLTDNGTILF	SQNVSNEANN	NGGAITTKTL	SISGNTSSTT	ETSNSAKKI.C
	201	GAIYSSAAAS	ISGNTGQLVF	MNNKGETGGG	ALGFEASSST	TONSSI, PECC
	251	NTATDAAGKG	GAIYCEKTGE	TPTLTISGNK	SLTFAENSSV	TOGGATCAHG
10	301	LDLSAAGPTL	FSNNRCGNTA	AGKGGAIAIA	DSGSLSLSAN	OGDITELGNE
10	351	LTSTSAPTST	RNAIYLGSSA	KITNLRAAOG	OSTYFYDDTA	SMTTCA SDUT.
	401	TINGPOSNSP	LDYSGTIVFS	GEKLSADEAK	AADNETSTLK	OPTALASOTT.
	451	ALKGNVELDV	NGFTQTEGST	LLMOPGTKLK	ADTEATSLTK	LAMMI, CAILED
	501	NKSVSIETAG	ANKTITLTSP	LVFODSSGNF	YESHTINOAF	TODIAMETAA
15	551	TAASDIYIDA	LLTSPVQTPE	PHYGYOGHWE	ATWADTSTAK	SCTMTSTUTTO
15	601	YNPNPERRAS	VVPDSLWASF	TDIRTLOOIM	TSOANSTYOO	PGT.WA SCTAM
	651	FFHKDKSGTN	QAFRHKSYGY	IVGGSAEDFS	ENIFSVAFCO	T.FCKDKDT.FT
	. 701	VENTSHNYLA	SLYLQHRAFL	GGLPMPSFGS	ITDMLKDTPI.	TT.MACT.CVCV
20	751	TKNDMDTRYT	SYPEAOGSWT	NNSGALELGG	ST.AT.VT.DKEA	DEFOCVEDET.
	801	KFQAVYSRQQ	NFKESGAEAR	AFDDGDLVNC	SIPVGIRLER	TSEDEKNMER
	851	TOTALIGDAA	RKNPRSRTSL	MVSGASWTSL	CKNLAROAFT	ASAGSHLTLS
	901	PHVELSGEAA	YELRGSAHIY	NVDCGLRYSF	*	

.Pmp 9 (CPn0447)

One example of a Pmp 9 protein is set forth as SEQ ID NO^s 33 & 34 in WO 02/02606. {GenBank accession number: gi|4376731|gb|AAD18591.1| 'CPn0447'; SEQ ID NO: 26 below}.

SEQ ID No 26

30	_		•			
30	_1	MKSSLHWFLI	SSSLALPLSL	NFSAFAAVVE	INLGPTNSFS	GPGTYTPPAQ
	51	TINADGTIYN	LTGDVSITNA	GSPTALTASC	FKETTGNLSF	QGHGYQFLLQ
	101	NIDAGANCIE	TNTAANKLLS	FSGFSYLSLI	OTTNATTGTG	ATKSTCACST
	151	QSNYSCYFGQ	NFSNDNGGAL	QGSSISLSLN	PNLTFAKNKA	TORGGAT.Vem
25	· 201	GGITINNTLN	SASFSENTAA	NNGGAIYTEA	SSFTSSNKAT	CETMICITAT
35	. 251	SATGGAIYCS	STSAPKPVLT	LSDNGELNFI	GNTAITSGGA	
	301	GGPTLFKNNS	AIDTAAPLGG	ATATADSGST.	SLSALGGDIT	TITOMENTOS
	351	SSSQTTTRNS	INIGNTNAKI	VOLRASOGNT	IYFYDPITTS	TTANICDALM
	401	LNGPDLAGNP	AYOGTIVESG	EKLSEARAAR	ADNLKSTIQQ	TIMALISDALIN
40	451	LKSGVTLVAK	SESOSPGSTI.	I.MDA CTTI.ET	ADGITINNLV	PULLAGGQLS
40	501	KATLKATOAS	OTVTLSGSLS	LANDESCHAVE	DVSWNNPQVF	TWADSTIKETK
	551	ANIHITDLAA	DPLEKNPTHW	GVOCNWAT.CM	QEDTATKSKA	SCLITLIADDP
	601	NPNPERRGTL	VANTIWGGEV	DIDETOOLIN	TKVRQSQETR	ATLITWIKTGY
	651	FHKDSTKINK	GERHTSAGVV	PAYOTÓOTAN	NLITAAFCQL	GIWCEGISNF
	701	KNRASAYAAS	LHT.OHT.ATT.C	CONTITUASD	SESEQPVLFD	FGKDRDHFIN
45 .	751	TMKTVVTOAD	KCECCMANDC	CALLIAGOES	HTALSHEGLE	AQISYIYSKN
	801	ASYTHODSEK	EDMINITIAN CE	CALELASSLE	HTALSHEGLF	HAYFPFIKVE
	851	VTVVANVVDV	TONUTTIONS	DOGNITIONSA	PIGITFERFS	RNERASYEAT
	901	T.EVPONT.CME	TRECORGINA	NNTSWKTTGT	NLSRQAGIGR	AGIFYAFSPN
	301	LEVTSNLSME	TRGSSKSYNA	DLGGKFQF*		

50 Pmp 11 (CPn0451)

One example of a Pmp 11 protein is set forth as SEQ ID NO^s 115 & 116 in WO 02/02606. {GenBank accession number: gi|4376733|gb|AAD18593.1| 'CPn0451'; SEQ ID NO: 27 below).

55 SEQ ID No 27

	1	MKTSIPWVLV	SSVLAFSCHL	OSLANEELLS	PDDSFNGNTD	COTETERNO
	51	TTYSLTGDVF	FYEPGKGTPI.	SDSCEKOTED	MI.TELCHOUS	IMECRIPATO
	101	HAGAAASTTA	NKNLTESGES	LICEDSSDST	MAILTONGUS	LIFGFIDAGT
60	151	RKLVVAGNFS	TADGGATKGA	SELT TOTOTO	AT ECHRICAGE	SAGGVNLENI
	201	ARIANNTGYV	RELSNIASTS	CCALDDECMO	ALIF SINKSSST	KGGALATTAG
	251	AICNTKASGS	DELTTONIUM	GGWIDDEGIS	TESNIKETAE	EGNAAKTTGG
	. 251	AICNTKASGS	PELIISNNKT	LIFASNVAET	SGGATHAKKI.	EGNAAKTTGG ALSSCGETER

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301 LRNNVSSATP KGGAISIDAS GELSLSAETG NITFVRNTLT TTGSTDTPKR
                     NAINIGSNGK FTELRAAKNH TIFFYDPITS EGTSSDVLKI NNGSAGALNP
                351
                     YQGTILFSGE TLTADELKVA DNLKSSFTQP VSLSGGKLLL QKGVTLESTS
                401
                451 FSQEAGSLLG MDSGTTLSTT AGSITITNLG INVDSLGLKQ PVSLTAKGAS
 5
                     NKVIVSGKLN LIDIEGNIYE SHMFSHDQLF SLLKITVDAD VDTNVDISSL
                501
                     IPVPAEDPNS EYGFQGQWNV NWTTDTATNT KEATATWTKT GFVPSPERKS
                551
                601 ALVCNTLWGV FTDIRSLQQL VEIGATGMEH KQGFWVSSMT NFLHKTGDEN
                     RKGFRHTSGG YVIGGSAHTP KDDLFTFAFC HLFARDKDCF IAHNNSRTYG
                651
                     GTLFFKHSHT LQPQNYLRLG RAKFSESAIE KFPREIPLAL DVQVSFSHSD
                701
10
                     NRMETHYTSL PESEGSWSNE CIAGGIGLDL PFVLSNPHPL FKTFIPQMKV
                751
                801
                     EMVYVSQNSF FESSSDGRGF SIGRLLNLSI PVGAKFVQGD IGDSYTYDLS
                851 GFFVSDVYRN NPQSTATLVM SPDSWKIRGG NLSRQAFLLR GSNNYVYNSN
901 CELFGHYAME LRGSSRNYNV DVGTKLDF*
                     CELFGHYAME LRGSSRNYNV DVGTKLRF*
15
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Pmp 12 (CPn0452)

One example of a Pmp 12 protein is set forth as SEQ ID NO^s 51 & 52 in WO 02/02606. {GenBank accession number: gi|4376735|gb|AAD18594.1 'CPn0452'; SEQ ID NO: 28 below).

20

SEQ ID No 28

	1	MTILRNFLTC	SALFLALPAA	AQVVYLHESD	GYNGAINNKS	LEPKITCYPE
25	51	GISTIFFDDA	RISNVKHDQE	DAGVFINRSG	NIFFMONDON	ETEUNII MTEC
25	. 101	FGAALSNRVG	DTTLTLSNFS	YLAFTSAPLL	POGOGATVST.	COMMITTENEDE
	151	VTFCGNYSSW	SGAAIYTPYL	LGSKASRPSV	MI.SCMDVIAE	RDNVSQGYGG
	201	AISTHNLTLT	TRGPSCFENN	HAVHDUNISNO	CATATARCCO	TOTOUTGE
	251	IFKGNTASQD	GNTTHNSTHI.	OSCAOEKNIT.D	MCECOLUE	TSTSVKSGDL
	301	ITDLVINAPE	GKETYEGTIS	ESCI.CL DDUR	MVSESGVIFI	LQDVTLAGGT
30	351	LSTSDGVTTO	LUCEVODACC	T CLUDDIE	VCAENLISTI	LQDVTLAGGT
	401	LSLSDGVTLQ	THOU TOT THE	TLIMSPGTTL	LCSGDARVQN	LHILIEDTON
	451	FVPVRIRAED	KDALIVSLEKL	KVAFEAYWSV	YDFPQFKEAF	TIPLLELLGP
		SFDSLLLGET	TLERTQVTTE	NDAVRGFWSL	SWEEYPPSLD	KDRRITPTKK
	501	TVFLTWNPEI	TSTP*			

35 Pmp 13 (CPn0453)

One example of a Pmp 13 protein is set forth as SEQ ID NO^s 3 & 4 in WO 02/02606. {GenBank accession number: gi|4376736|gb|AAD18595.1 'CPn0453'; SEQ ID NO: 29 below}.

40 SEQ ID No 29

	1 MKTSIRKFLI	STTLAPCFAS	TAFTVEVIMP	SENFDGSSGK	TEPYTTI.SDD
	SI RGTLCIFSGD	LYIANLDNAI	SRTSSSCFSN	RAGALOTICK	COVEREINTE
45	01 SSADGAAISS	VITQNPELCP	LSFSGFSQMI	FONCESLTSD	TEACHTINIA
45	51 SAIYATTPML	FTNNDSILFO	YNRSAGFGAA	TROTSTTTEM	TUVOLITIM
20	01 GSISNGGALT	GSAAINLINN	SAPVIFSTNA	TCTVCCATVI	TOCOME
25	51 LSGVLFVNNS	SRSGGAIYAN	GNVTFSNNSD	T.TEONNEY CD	OMOTERATION
30	01 PTPPAVTPLL	GYGGATECTP	PATPPPTGVS	TOTOGONOUS	QNSLIPAPTPP
35	51 GALYGKKIST	DSNKSTTELG	NTAGKGGAIA	TITOGENOVI	FLENIASEQG
50	1 NLSITSGTPT	DNSTURCEDA	MINGROGATA	IPESGELSLS	ANQGDILFNK
45		ADCAVCCTTV	KFATLGATQG	ATPARADBIL	SDDLSAASAA
50	ון מינ. און פאני איני איני	MUDICACOURT	FSGETLTATE	AATPANATST	LNQKLELEGG
55	TAIL DOLD DOWN	NVHNFTQDEK	SVVIMDAGTT	LATTNGANNT	DGAITLNKLV
.60	TNEDSEDGIK	AAVVNVQSTN	GALTISGTLG	LVKNSQDCCD	NHGMFNKDLQ
EE .	OVPILELKAT	SNIVITIDES	LGTNGYQQSP	YGYQGTWEFT	IDTTTHTVTG
•••	T MAKKIGATEH	PERLAPLIPN	SLWANVIDLR	AVSOASAADG	POWDOWOT OF
70	I IGIINFFHAN	HTGDARSYRH	MGGGYLINTY	TRITPDAALS	LCECOLETVO
75	T KDATAGHCHS	NVYFATVYSN	ITKSLFGSSR	FFSGGTSRVT	VSRSNEKVKT
80	1 SYTKLPKGRC	SWSNNCWLGE	LEGNLPITLS	SRILNIKOTT	DEMKYEMYAY
85	1 THGGIQENTP	EGRIFGHGHL	LNVAVPVGVR	FCKNSHNDDD	PUTTURVALA
60 90	1 DVYRHNPDCD	TTLPINGATW	TSIGNNLTRS	TLLMORECUT	CIMIDIA ELEC
95	1 HCGCDIRRTS	ROYTLDIGSK	T.RF*	TTT A AUGGUT	PANDAPETEG

Pmp 14 (CPn0454)

One example of a Pmp 14 protein is set forth as SEQ ID NOs 35 & 36 in WO 02/02606. {GenBank accession number: gi|4376737|gb|AAD18596.1 'CPn0454'; SEQ ID NO: 30 below}.

SEQ ID No 30

5

	1	MPLSFKSSSF	CLLACLCSAS	CAFAETRLGG	NFVPPITNOG	EEILLTSDFV
10	51	CSNFLGASFS	SSFINSSSNL	SLLGKGLSLT	FTSCOAPTNS	NYALLSAAET
10	101	LTFKNFSSIN	FTGNQSTGLG	GLIYGKDIVF	OSIKDLIFTT	MRWAVCDACW
	151	TTSATPAITT	VTTGASALOP	TDSLTVENIS	OSIKEFGNI.A	NEGGATGGGD
	201	TAVVKFINNT	ATMSFSHNFT	SSGGGVIYGG	SSLLFENNSG	CTTETANGCU
•	251	NSLKGVTPSS	GTYALGSGGA	ICIPTGTFEL	KNNOCKCTES	VAICHDAIDA CA
45	301	IYAETCNIVG	NOGALLLDSN	TAARNGGAIC	AKVINTOGPG	DIFFEDNDAGA
15	351	KGGAIFIGPS	VGDPAKOTST	LTILASEGDI	AFOCHMINTE	DOTONATORO
	401	AGGEIVSLSA	OGGSRLVFYD	PITHSLPTTS	DOMEDITATION	PGTKWATIVE
	451	SKGLSSTELL	LPANTTTILL	GTVKIASGEL	KITDMATAMA	I CENTOCOCO
	501	LTLGSGGTLG	LATPTGAPAA	VDFTIGKLAF	MITIDIAY VIV	TIGHATQGSGQ
	551	NVTLTGALVL	DEHDVTDLVD	MVSLQTPVAI	DIAUEVCAMI	VSASVNAGTK
20	601	ATPSHYGYOG	KWSYTWSRPI	LIPAPDGGFP	CCDCDCYMMI	TRIGEPOGEL
	651	RSTYLLDPER	YGETVSNSLW	ISFLGNQAFS	DILONALTE	YAVWNSDTLV
	701	LGAYVEHTDR	UCHECECCDA	GGYQAALSMN	DITIONALITY	HPGLSITAKA
	751	NPYDSPCSEO	MVILEPPOOR	PIVTQKSEAL	TOHTTLGLS	FGQLYGKTNA
	801	BDKY BK60G0	MUNNOVONT	PIVIQASEAL	ISWKAAYGYS	KNHLNTTYLR
25		T.CCMOCKEAR	MUMARITATI	SAEHPFLNWC	LLTRPLAQAW	DLSGFISAEF
	901	T.VANDUANTABA	MOUNTAINMEN	GKGYNVSLPI	GCSSQWFTPF	KKAPSTLTIK
	951	TOVETTENTIKA	GKNGFTNHRV	NQESTSISGA	NLRRHGLFVQ	IHDVVDLTED
	221	TOWLTMITTED	GKNGFTNHRV	STOLKSTER		

30 Pmp 15 (CPn0466)

One example of a Pmp 15 protein is set forth as SEQ ID NO^s 5 & 6 in WO 02/02606. {GenBank accession number: gi|4376751|gb|AAD18608.1 'CPn0466'; SEQ ID NO: 31 below}.

35 SEQ ID No 31

	1	MRFFCFGMLL	PFTFVLANEG	LQLPLETYIT	LSPEYOAAPO	VGFTHNQNQD
	51	LAIVGNHNDF	ILDYKYYRSN	GGALTCKNLL	ISENIGNVFF	EKNYCDNSCC
40	101	AIYAAQNCTI	SKNQNYAFTT	NLVSDNPTAT	AGSLLGGALF	ATNOSTOWN.
40	. 151	GQGTFVDNLA	LNKGGALYTE	TNLSIKDNKG	PITIKONRAL.	NSDSI.CCCTV
	201	SGNSLNIEGN	SGAIQITSNS	SGSGGGIFST	OTUTTSSNKK	I.TETCENICAE
	251	ANNYGSNFNP	GGGGLTTTFC	TILNNREGVI.	FNNNOSOSNG	CATUAVOTTT
	301	KENGPVYFLN	NTATRGGALL	NLSAGSGNGS	FILSADNODI	GUTTIMUSTIT
4.5	351	ALNPPYRNAI	HSTPNMNLQI	GARPGYRVI.E	VDDTEUET.DC	CEDIT ENERGY
45	401	GHTGTVLFSG	EHVHQNFTDE	MNFFSVI.DNT	SELBOOM VA	SPEIDINFET
	451	FFORGGTLLL	GQGAVITTAG	TTDTDCCTDT	THE TRUE OF THE STREET	EDGAGLACYK
	501	FOAOAPKIWI	YPTKTGSTYT	EDGNDTTTTC	GTLTLRNSNN	TAIDLPSILS
	551		IVDVAAOKIN		CODINGNO	EDPYDSLDLS
	601		NTKHKLLYAN	~	SGEHYGYQGI	WSTYWVETTT
50	651		EKGHAASLQG		ERRGEFITNA	LWQSAYTALA
	701	CCOCDATION O			NGFKGFRSHM	TGYSATTEAT
	751		FAQFFSKAKE	HESQNSTSSH	HYFSGMCIEN	TLFKEWIRLS
			HTHTMYQGLL	EGNSQGSFHN	HTLAGALSCV	FLPQPHGESL
	801	OTARELIATY	IRGNLAAFQE	SGDHAREFSL	HRPLTDVSLP	VGIRASWKNH
55	851	HRVPLVWLTE	ISYRSTLYRQ	DPELHSKLLI	SOGTWITOAT	PVTYNALGIK
55	901	VKNTMQVFPK	VTLSLDYSAD	ISSSTLSHYL	NVASRMRF*	

Pmp 16 (CPn0467)

One example of a Pmp 16 protein is set forth as SEQ ID NO^s 7 & 8 in WO 02/02606. {GenBank accession number: gi|4376752|gb|AAD18609.1| 'CPn0467'; SEQ ID NO: 32 below}.

SEQ ID No 32

	1	MFGMTPAVYS	LQTDSLEKFA	LERDEEFRTS	FPLLDSLSTL	TGFSPITTFV
5	51	GNRHNSSQDI	VLSNYKSIDN	ILLLWTSAGG	AVSCNNFLLS	MAEDHVEECK
3	101	NLAIGTGGAI	ACQGACTITK	NRGPLIFFSN	RGLNNASTGG	ETRECATACM
	151	GDFTISQNQG	TFYFVNNSVN	NWGGALSTNG	HCRIOSNRAP	S.T.FEMMTA DC
	201	GGGALRSENT	TISDNTRPIY	FKNNCGNNGG	AIOTSVTVAT	KNNSCSVIEN
	251	NNTALSGSIN	SGNGSGGAIY	TTNLSIDDNP	GTILFNNNYC	TRRCCATOTO
10	301	FLTIKNSGHV	YFTNNQGNWG	GALMLLODST	CLLFAEOGNT	A FONNEYET.T
	351	TFGRYNAIHC	TPNSNLQLGA	NKGYTTAFFD	PIEHOHPTTN	PLTENDNANH
	401	QGTILFSSAY	IPEASDYENN	FISSSKNTSE	LRNGVISTED	RAGMOEVKET
	451	QKGGILKLGH	AASIATTANS	ETPSTSVGSO	VIINNIATNI.	PSTLAKCKAD
	501	TLWIRPLQSS	APFTEDNNPT	ITLSGPLTLL	NEENRDPYDS	TDLSEDLONT
45	551	HLLSLSDVTA	RHINTDNFHP	ESLNATEHYG	YOGIWSPYWV	ETTTTTTMING
15	601	IETANTLYRA	LYANWTPLGY	KVNPEYOGDL	ATTPLWOSFH	TMFSLLDSVN
	651	RTGDSDIERP	FLEIQGIADG	LFVHONSIPG	APGFRIOSTG	VSLOASSETS
	701	LHQKISLGFA	QFFTRTKEIG	SSNNVSAHNT	VSSI-VVEL DW	FORAFATOTU
20	751	LAYGYGDHHL	HSLHPSHQEO	AEGTCYSHTL	AAATGCSFPW	COKSVI.HI.CD
	801	FVQATAIRSH	QTAFEEIGDN	PRKFVSOKPF	YNLTLPLGTO	CKMUCKERIAD
	821	TEWTLELSYO	PALAOONBOI	GVTLLASGGS	WDILGHNYVR	NALGYKVHNO
	901	TALFRSLDLF	LDYQGSVSSS	TSTHHLQAGS	TLKF*	

Pmp 18 (CPn0471)

One example of a Pmp 18 protein is set forth as SEQ ID No 33 below {GenBank accession number: gi|4376753|gb|AAD18610.1| 'CPn0471'.

SEQ ID No 33

30	_						
30	. 1	MQNNRSLSKS	SFFVGALILG	KTTILLNATP	LSDYFDNQAN	QLTTLFPLID	TLTNMTPYSH
	9.1	KATLFGVRDD	INODIVLDHO	NSIESWFENF	SODECALSER	CT. A TURNITURATO	TT DT MOUNTYS
	727	RAGAMYVNGN	FDLSENHGSI	IFSGNLSFPN	ASNEADTOTO	CAMILICENTUM	TOWNOOMAND
	707	TNNKAKSSGG	AIQAAIINIK	DNTGPCLFFN	NAAGGTAGGA	T. FANTA COTENT	MCODIVETIME
25	241	GOGLGGATKA	HQECILTKNT	GSVIFNNNFA	MEADICANUC	SCCATVOTOO	CTIMITOGTAN
35	301	FUNNTAARDG	GAICTQSLTI	ODSGPVYFTN	NOGTWCCATM	T.PODGA CTIT.E	ADOCDETENAL
	201	MRHFKDTFSN	HVSVNCTRNV	SLTVGASOGH	SATEVDDILO	DVTTOMETOV	TENTENT CO.
	421	TULSSIAIDD	TSTSRDDFIS	HFRNHIGLYN	·GTLALEDRAE	WKWVKEDOEC	CULDI CODAM
	401	FSTIDEEQSS	SSVGSVININ	NLAINLPSIL	GNRVAPKT.WT	PPTCCCADVC	EDMINTER O
	541	GPLSLLDDEN	LDPYDTADLA	OPIAEVPLLY	LIDVTAKHTM	TDNFYPEGLN	EDMMETIMES
40	601	WSPYWIETIT	TSDTSSEDTV	NTLHROLYGD	WTDTGVKIMID	ENKGDIALSA	TTOHYGYOGV
	661	TLRYOTOOGO	IAPTASGEAT	RIFVHONSMN	DAKGERMEYE	GYSLGTTSNT	FWQSFHNLFA
	721	SOLFSNLYES	HSDNSVASHT	TTVALATION	DAKGEMIEAT	SLAYSYSNHH	ASNHSFGVNF
	781	OTEGKCYSTT	T.GAAT.SCST.S	TOWNSHIP	WINDERPSISA	SNQTAFQESG	IKASGYSGKI
	841	PLYNTTVPLG	TOSAMESKED	LUTANAMENT	1PF1QATAVR	SNOTAFQESG EINVSLESSG	DKARKFSVHK
45	901	ARNAIAFKGR	NOTETEDATE	DELIMINTERY	TOPATTOOMS	EINVSLESSG	SSWLLSGTTL
•	302		MATETERITO	ATTINITY GRAZE	SSTTTHYLHA	GTTFKF	

Pmp 19 (CPn0539)

One example of a Pmp 19 protein is set forth as SEQ ID No 34 below {GenBank accession number: gi|4376829|gb|AAD18679.1 'CPn0539'; SEQ ID NO: 34 below}.

SEQ ID No 34

	1	MKQMRLWGFL	FLSSFCQVSY	LRANDVLLPL	SGIHSGEDLE	דמ פפפק נודק נו	KTTYSLRKDF
	91	IVCDFAGNSI	HKPGAAFLNL	KGDLFFINST	PLAALTERNT	HI CAPCACI P	CECKEREN
55	141	HSLVLENNES	WGGVLTTSGD	LSFINNTSVL	CONNISYCEC	CALLLOCAVO	VAT BEDDAMA
	TOT	TTTLTKNKVA	NODESHPGYG	GAVSSTSPGS	PTTFADMORT	LEORNEOUTA	CA TIDMOGRA
	241	TREMMIQITS	FFSNKASFGG	AVYSRYCNLY	SOWGDTLETK	እየአ አ አ አሚያረግር አ ተ	TIA DIGITITADO
	201	VC2TALEEN2	ATAGGAIAVN	AVCDINAGGP	VRFINNSALC	LNICCATUMOA	MCOTT DT WAR
00	201	QGDIEFCGNK	VRSOFHSHIN	STSNFTNNAT	TTOGADDREE	LCANDOUDTO	TIPDTTO
60	421	MINZPITINHO	RLLEAGGAVI	FSGARLSPEH	KKENKNKTCT	TMODURE CCC	TH GIRGON TY
	401	AVKSFIQEGG	PTATGEGREE	TTOGKNSEKD	KIVITNLGFN	LEMILDOCDDA	PIDAMERAGE
	541	EISGVPRVYG	HTESFYENHE	YASKPYTTSI	ILSAKKLVTA	PSPDEKDION	I.TTA DODUMO

	601 YGYQGSWEFS	WSPNDTKEKK	TIIASWTPTG	EFSLDPKRRG	SETDOWNINGO	ECOLMEN ONE
	OOT ANMATHURE	VIPLOHLCVF	GGPVYOIMEO	NPKOSSNNT.T.	VOHACHMUCA	DIDECEMENT
	721 SAALTQLESS	SSOONVADKS	HAOILIGTVS	LNKSWOAT.St.	Deepeymeno	017447777777777777777777777777777777777
5	OT GISKGSMKNY	GWSGSVGMSY	AYPKGIRYLK	MTPFVDIAVT	KT.UOMDERREE	Olmphureee
3	OAT EMINDSPATE	IALEMRFIGS	RSSLFLQVST	SYIKDLRRVN	POSSASTATA	HYTWDIOGVP
	901 LGKEALNITL	NSTIKYKIVT	AYMGISSTQR	EGSNLSANAH	ACTISTISE	

Pmp 20 (CPn0540)

One example of a Pmp 20 protein is set forth as SEQ ID NO^s 119 & 120 in WO 02/02606. {GenBank accession number: gi|4376830|gb|AAD18680.1 'CPn0540'; SEQ ID NO: 35 below}.

SEQ ID No 35

15						
15	1		AAVLPALTAI	GDPASVEIST	SHTGSGDPTS	DAALTGFTOS
	51	STEIDGITYI	: IVGDITFSTE	TNIPVPVVTP	DANDSSSNSS	KCCCCCCCXT
	101	STIKSSNTHS	6 DFDFTKDSVI	DLYHLFFPSA	SNTT-NPAT-T-S	9999000000
	151	SSSSSSSSAS	AVVAADPKGO	AAFYSNEANG	TITETTOSCA	DCGT.TT.ONT.W
20	201	MIGDGAAIYS	KGPLVFTGLE	NLTFTGNESO	KSGGAAYTEG	ል፤.ጥጥ ስል ፕሬነር፣አ
20	251	VIFIGNISAG	QGGAIYVKEA	TLFNALDSLK	FERNTSGOAG	CCTVTECTT
	301	TSNITKSTER	' ISNKASVPAP	APEPTSPAPS	SLINSTITIOT	STLOTEDANCE
	351	IPAVAPVAAV	TPTPISTQET	' AGNGGAIYAK	OGISISTEKD	T.TEKSNISA CW
	401	DATLIVDSSI	' IGESGGAIFA	ADSIOIOOCT	GTTLFSGNTA	MKSCCCTVAN
25	451	GOVILEDIAN	LKMTNNTCKG	EGGAIYTKKA	LTTNNGATI.T	TECCMITCHINA
23	501	GGATFAVGGT	TLSDLVEVRF	'SKNKTGNYSA	PTTKAASNTA	DITTICC COTON N
	551	SPAVPAAAAA	PVINAAKGGA	LYSTEGLTVS	GITSTLEFEN	NECOMOCCON
	601	IVIKIFQCSD	SHRLQFTSNK	AADEGGGLYC	CDDVTT.TMLT	CKTT.POPMEC
	651	EXHGGGLSLA	SGKSLTMTSL	ESFCLNANTA	KENGGGAMID	ENTITE TENTO
30	701	PTPNEPAPVQ	QPVYGEALVT	GNTATKSGGG	IYTKNAAFSN	T.SSUTEDONT
30	751	SSENGGALLT	QKAADKTDCS	FTYITNVNIT	NNTATCNCCC	TACCYAURDD
	801	TDNLTAGSNG	AKKGGGVYLE	DALILEKVIT	GSVSONTATE	SCCCTVAVDT
	. 851	QLQALPGSFT	ITDNKVETSL	TTSTNLYGGG	TVSSCAUTT.T	MTCCTTCCTTCC
	901	NSVINTATSQ	DADIOGGGIY	ATTSLSTNOC	NTDIT. FCMMC	$\lambda \lambda m \nu \nu m c m m \nu$
35	951	QIAGGAIFSA	AALTENNSOD	IIFLNNSAKS	EATTAATACM	KDCCCCATAA
33	1001	MOATPIMMEE	TTFKGNYAET	GGAIGCIDIT	NGSDDDKNGT	ADMOCUT BOD
	1051	NSALNRGGAL	YGETIDISRT	GATFIGNSSK	HDGSATCCST	AT.TI.A DMCOT
	1101	TEENNKATET	TATTKASINN	LGAAIYGNNE	TSDVTTSLSA	PMCCTPPVM
	1151	LCTAINKYCS	IAGNVKFTAI	EASAGKAISF	YDAVNUSTKE	THIA OFT. ET. NE
40	1201	KATSTGTILF	SGELHENKSY	IPOKVTFAHG	NLILGKNAET.	SWISETORDO
40	1251	TTTTMGPGSV	LSNHSKEAGG	IAINNVIIDE	SETUPTEDNA	יז זע.זייםם מזיי
	1301	SKINADSKOK	IDITGTVTLL	DPNGNLYONS	YLGEDROTTI.	FNITINICACCA
	1351	ATMALLÓG	NLGAKKGYLG	TWNLDPNSSG	SKITLKWTED	KVT.DWDVTDD
	1401	DMHLAIMSIM	GAQNSLVTVK	OGILGNMLNN	ARFEDDARM	TWACA TOOPT
45	1451	KKEVSKNSDS	FTYHGRGYTA	AVDAKPROEF	TIGAARSOVE	CHAPCEVIII
40	1501	NYKHKGSGHS	TQASLYAGNI	FYFPAIRSRP	TIFOGVATVG	VMOUDTTTTV
	1551	POTEEKNMAN	WDSIAWLFDL	RFSVDLKEPO	PHSTARLTEV	TEAEVTOIDA
	.1601	EKLIETDADD	RSFSACSYGN	LAIPTGFSVD	GATAWPETTI.	VATETICA A UT D
	1651	ATTKNN5KVL	YEVLSTKEKG	NVVNVLPTRN	AARAEVSSOI	YLGSYWTLYG
	1701	TYTIDASMNT	LVQMANGGIR	FVF*		
50					•	

Pmp21 (CPn0963)

One example of a Pmp 21 protein is set forth as SEQ ID NO^s 83 & 84 in WO 02/02606. {GenBank accession number: gi|4377287|gb|AAD19099.1| 'CPn0963'; SEQ ID NO: 36 below}.

SEQ ID No 36

	1	MVAKKTVRSY	RSSFSHSVIV	AILSAGIAFE	AHSLHSSELD	LGVENKOFEE
60	27.	HSAHVEEAQT	SVLKGSDPVN	PSOKESEKVL	YTOVPLTOGS	SCESTIDIADA
00	101	MEDEHEGHUE	EETTVFGIDQ	KLVWSDLDTR	NESOPTOEPD	TSNAVSEKTS
	121	SUTKENRKUL	ETEDPSKKSG	LKEVSSDLPK	SPETAVAAIS	EDIETSENTS
	201	ARDPLQGLAF	FYKNTSSOSI	SEKDSSFOGT	TESGSGANSG	T.CEPNILVA DV

		251	SGAAVYSDRI	IVFENLVKGL	SFISCESLED	GSAAGVNIVV	THCGDVTT.TD
		301	CWIGNDLEAL	RLVKDFSRGG	AVFTARNHEV	OMNTACCTIC	*********
		351	. EVMONEKSNG	GAFACGSFVY	SNNENTALWK	ENOAL SCOAT	COLOREDTOS
5		401	MCSWIELSCN	OSLIALGEHI	GLTDFVGGGA	T A A COUNT OF THE	ATTA THE COURT
Э		451	TOWINGGAIL	AGTVDLNETI	SEVAFKONTA	ALTCCALCANT	DVUTTANDER
		501	EILFEQNEVR	NHGGAIYCGC	RSNPKLEQKD	SCENINITION	DAVITANNEG
		551	ASVLEVMTOA	EDYAGGGALW	CHNVLLDSNS	CNICETONIC	SGATTFLKNK
		601	GGGAILSTDR	VTTSNNSGDV	VFKGNKGQCL	CUITOLIGNIC	GSTFWIGEYV
		651	NKDEKSLNAC	SHGDHVDDKT	VEEEVPPSLL	ACKIVAPOET	APVESDASST
10		701	HIFITDNTGN	LPESCHLOCO	EESSTVGDLA	EEHPVVSSTD	IRGGGAILAQ
		751	VVFSDNVTSN	CCDCCCATTA	EESSI VGDLA	IVGGGALLST	NEVNVCSNON
		801	SUNTEDNICEN	GCDSGGWTTW	KKVDISANHS	VEFVSNGSGK	FGGAVCALNE
		851	SENOPSCCCX	VSFSKMKIKL	GGAGVAAPQG	SVTICGNQGN	IAFKENFVFG
		901	CHICAGOGA CHICAGOGA	TIAMSSANIO	DNAGDILFVS	NSTGSYGGAI	FVGSLVASEG
15		951	CMICEACHDS	NSGDILLFAKN	STQTAASLSE	KDSFGGGAIY	TQNLKIVKNA
		1001	CONDCUTURE	PSGAGVQIAD	GGTVCLEAFG	GDILFEGNIN	FDGSFNAIHL
		1051	CGMDSKIAEP	SAVQDKNIIF	QDAITYEENT	IRGLPDKDVS	PLSAPSLIFN
		1101	SYLOUDSHOH	HEGTIRFSRG	VSKIPOTAAT	OFCITATION.	A FOT BUT A CIT TOO
	•		PIGSSIATRY	GSTTKILDSO	VDSSAPT.PTF	NKEETLUCAC	MOTATIOCODED
20		1151	WOWNADIBA	LADITSITVD	LSSEVPEODG	דדדםםם.זס.זיוי	DECREE HOLES
20		1201	TDTKTTDBJM	VGYENHALLS	SHKDIPLIST.	KTARCMTCTD	TADACT ONTER
		1251	TDASTEDITE	ATYGHTGVWS	ESKMEDGRIJ	VCWODTCVVI.	MDEVOCATAR
		1301	MMINGMITT	RALKOEIFAH	HTTAORMELD	FCTMMMCCCT	CHREDONITA
		1351	PL DG L VHHT.I.	GYALGLDTOL	VEDFLIGGGE	SOFFCKTECO	CVVXVXTXXXX
25		1401	THOMMINGILL	AGPWLIKGAF	VYGNINNDLT	ヤカマです。 こすぐか	CCMTCVCDTA
25		1451	GISIDIKITA	NPRREISAIV	STVVPFVEAE	VVD TOT. DETC	POCKERMON
	•	T 2 O T	WIKE EM AWID	FGFALEHAYS	RGSRAEVNSV	OLYMPHIA COLUMN	VCDUCT TOT TO
		1551	DAAYSWKSYG	VDIPCKAWKA	RISNNTEWNS	ATTAINEDAIK	WAS ASPILIFY
		1601	FNGGIRIIF*			THOTTHEMY	EMKEDLTAAD

Preferred PMP proteins for use with the invention comprise an amino acid sequence: 30 (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to one of the polypeptide sequences set forth for the pmp proteins above and/or (b) which is a fragment of at least n consecutive amino acids of one of the polypeptide sequences set forth above wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 35 60, 70, 80, 90, 100, 150, 200, 250 or more). These PMP proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of the polypeptide sequences set forth above. Preferred fragments of (b) comprise an epitope from one of the polypeptide sequences set forth above. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-40 terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of one of the polypeptide sequences set forth above. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular 45 domain).

Fifth Antigen Group

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The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group or the third antigen group or the fourth antigen group. Such other *Chlamydia pneumoniae* antigens include a fifth antigen group consisting one or more cell surface exposed proteins. These antigens are referred to herein as the "fifth antigen group". Each of the *Chlamydia pneumoniae* antigens of the fifth antigen group is described in more detail below.

(37) PorB Outer Membrane Protein B (CPn0854)

One example of a PorB protein is set forth as SEQ ID NO⁵: 67 & 68 in WO 02/02606. {GenBank accession number: gi|4377170|gb|AAD18992.1| 'CPn0854'; SEQ ID NO: 37 below}. Preferred PorB proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 37; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 37, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These PorB proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 37. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 37. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 37. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 37

5

10

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20						
20		1 <u>MNSKMLKHLR</u>	LATLSFSMFF	GIVSSPAVYA	LGAGNPAAPV	LPGVNPEQTG
	5	1 WCAFQLCNSY	DLFAALAGSL	KFGFYGDYVF	SESAHITNVP	VITSVTTSGT
	10	1 GTTPTITSTT	KNVDFDLNNS	SISSSCVFAT	IALOETSPAA	IPLLDIAFTA
	15	1 RVGGLKQYYR	LPLNAYRDFT	SNPLNAESEV	TDGLIEVOSD	YGIVWGLSLO
05	20	1 KVLWKDGVSF	VGVSADYRHG	SSPINYIIVY	NKANPEIYFD	ATDGNLSYKE
25	25	l WSASIGISTY	LNDYVLPYAS	VSIGNTSRKA	PSDSFTELEK	OFTNFKFKIR
	. 30	1 KITNFDRVNF	CFGTTCCISN	NFYYSVEGRW	GYQRAINITS	GLOF*

(38) 76kDa Protein Homolog (CPn0728)

One example of a 76kDa Protein Homolog protein is set forth as SEQ ID NOs: 13 & 30 14 in WO 02/02606. {GenBank accession number: gi|4377033|gb|AAD18867.1| 'CPn0728'; SEQ ID NO: 38 below}. Preferred 76kDa proteins homologs for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 38; and/or (b) which is a fragment of at 35 least n consecutive amino acids of SEQ ID NO: 21, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These 76kDa protein homologs include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 38. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 38. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 38. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 38

	1	MVNPIGPGPI	DETERTPPAD	LSAQGLEASA	ANKSAEAORI	AGAEAKPKES
	51	KTDSVERWSI	LRSAVNALMS	LADKLGIASS	NSSSSTSRSA	DVDSTTATAP
50	101	TPPPPTFDDY	KTQAQTAYDT	IFTSTSLADI	QAALVSLODA	VTNIKDTAAT
	151	DEETAIAAEW	ETKNADAVKV	GAQITELAKY	ASDNOALLDS	LGKLTSFDLL
	201	QAALLQSVAN	NNKAAELLKE	MQDNPVVPGK	TPAIAOSLVD	QTDATATQIE
	251	KDGNAIRDAY	FAGQNASGAV	ENAKSNNSIS	NIDSAKAAIA	TAKTQIAEAQ
	301	KKFPDSPILQ	EAEQMVIQAE	KDLKNIKPAD	GSDVPNPGTT	VGGSKOOGSS

5	451 501 551 601	GVPPAAASSI NDATRDVINN VYSQVSALQS STQKFIAKLE	AAAALADAQK GSSVKQLYKT VSTPALTRSV VMQIIQSNPO	ALEAALGKAG SKSTGSDYKT PRARTEARGP ANNEEIROKL	QQQGILNALG QISAGYDAYK EKTDQALARV TSAVTKPPOF	AAQQELAAQA QIASAAVVSA SINDAYGRAR ISGNSRTLGD GYPYVQLSND NIGSLYSGYL
	651	Q*	SUPAEGSRTA	AEIKALSFET	NSLFIQQVLV	NIGSLYSGYL

(39) OmpA conserved outer membrane protein (CPn0695)

One example of an OmpA conserved outer membrane protein protein is set forth as 10 SEQ ID NOs: 59 & 60 in WO 02/02606. {GenBank accession number: gi|4376998|gb|AAD18834.1| 'CPn0695'; SEQ ID NO: 39 below}. Preferred ompA proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 15 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 39; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 39, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 39. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 39. Other preferred fragments lack one 20 or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 39. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of 25 a transmembrane domain, or of an extracellular domain).

SEQ ID No 39

20	1	MKKLLKSALL	SAAFAGSVGS	LOALPVGNPS	DPSLITTICTT	WEGAAGDPCD
30	51	PCATWCDAIS	LRAGFYGDYV	FDRILKVDAP	KTESMCAKOT	GSAAANYTTA
	101	VDRPNPAYNK	HLHDAEWFTN	AGFIAINTWD	PEDVECTICA	SNGYIRGNST
	151	AFNLVGLFGV	KGTTVNANEL	PNVSLSNGVV	FLVTDTCHGA	SVGARGALWE
	201	CGCATLGAEF	QYAOSKPKVE	ELNVICAVSO	EGIVIADACAA	GVAFPLPTDA
05	251	GVATATGTKS	ATINYHEWOV	GASLSVRLNS	LUDVICUOMO	RATFDADNIR
35	301	IAQPKLPTAV	LNLTAWNPSL	L'GNATAL STT	Describions	SCQINKFKSR
	351	KACGVTVGAT	LVDADKWSLT	AEARLINERA	AHVSGOFRE*	SCOTINGERSE

(40) PepA (CPn0385)

One example of a PepA protein protein is set forth as SEQ ID NOs: 99 & 100 in WO 02/02606. {GenBank accession number: gi|4376664|gb|AAD18529.1 'CPn0385'; 40 SEQ ID NO: 40 below}. Preferred PepA proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 40; and/or (b) which is a fragment of at least n consecutive amino 45 acids of SEQ ID NO: 40, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These PepA proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 40 Preferred fragments of (b) comprise an epitope from SEQ ID NO: 40. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, **50** 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 40. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 40

	1	MVLFHAQASG	RNRVKADAIV	LPFWHFKDAK	NAASFEAEFE	PSYLPALENF
_	51	QGKTGEIELL	YSSPKAKEKR	IVLLGLGKNE	ELTSDVVFOT	VATI.TRUIT.DV
5	101	AKCSTVNIIL	PTISELRLSA	EEFLVGLSSG	ILSLNYDYPR	YNKVDRNLET
	151	PLSKVTVIGI	VPKMADAIFR	KEAAIFEGVY	I TRDI VNEND	DEITPKKLAE
10	201	VALNLGKEFP	SIDTKVLGKD	AIAKEKMGI.I.	LAVSKGSCOD	PHFIVVRYOG
	251	RPKSKDHTVL	IGKGVTFDSG	GLDLKPGKSM	TAMKEDMAGG	ATVLGILSAL
	301	AVLELPINVT	GIIPATENAI	DGASYKMGDV	YVGMSGT.SVE	TOCHDYDON
	351	ILADAITYAL	KYCKPTRIID	FATITGAMIN	SIGEENACEE	SNNDVLAEDL
	401	LEASAETSEP	LWRLPLVKKY	DKTTHSDIAD	MUNITOCATIA	SMMDVLAEDL
	451	FLEESSVAWA	HLDTAGTAYH	EKEEDDADKA	ACCECIMANT	ATTAALFLOR
				HUBBRIERI	WORLGAKOIT	YYLENSLSK*

15 (41) Conserved Outer Membrane Protein (Cpn0278)

One example of a conserved outer membrane protein protein is set forth as SEQ ID NO: 41 below. GenBank Accession No. GI:4376552; AAD18427.1. conserved outer membrane proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 41; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 41, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These conserved outer membrane proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 41. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 41. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 41. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 41

	121 181	RLKSQKKLTI	AIPVDRTNAQ DVDAAVIPGN	RALHLLEECG FAIAANLSPK	RYDCKGELVV LIVCKGPANL KDSLCLEDLS	IAKVHLEPQA	GIKLKILPVD IYSKKHSSLE ENRSINILEV RSEDVGSPKM
40			OVQHEEDIKI	HGNITIMIQD	NG		•

Sixth Antigen Group

The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group or the third antigen group or the fourth antigen group or the fifth antigen group. Such other *Chlamydia pneumoniae* antigens include a sixth antigen group consisting one or more FACS positive CPn antigens. These antigens are referred to herein as the "sixth antigen group". Each of the *Chlamydia pneumoniae* antigens of the sixth antigen group is described in more detail below.

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(42) Predicted Omp (CPn0020)

One example of a predicted Omp protein is set forth as SEQ ID NO⁵: 91 & 92 in WO 02/02606. {GenBank accession number gi|4376272|gb|AAD18173.1: 'CPn0020'; SEQ ID NO: 42 below). Preferred Omp proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 42; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 42, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These Omp proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 42. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 42. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 42. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 42

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20	1	MKRCFLFLAS	FVLMGSSADA	LTHQEAVKKK	NSYLSHFKSV	SGIVTIEDGV
	51	THINNTKIQ	ANKVYVENTV	GOSLKLVAHG	NVMVNVDART	LACOUT DAME
	101	DTDSCLLTNG	RFAMYPWFLG	GSMITLTPET	TVTPKCVTCT	SECDANDI GI
	151	SGDYLEYSSD	SLLSIGKTTL	RVCRIPTLEL	PDESTMONET	DEGENADUCL
05	201	TGGFLGSYLG	MSYSPISRKH	FSSTFFI.DSF	PKHCACMCEN	FYLLING
25	251	NVFNMKSYYA	HRLAIDMAEA	HDRABI'HGDE	CETUVUTATIO	THCSOKOAPE
	301	TVADIFPNNF	MLKNTGPTRV	DOTWINDSTREE	CVI MCCIMANI	GEYHLSDSWE
	351	YLTLROYPIS	IYNTGVYLEN	TVECCVINES	GIDISOAKAN	SFQNANQELP
	401	LHKTVPLPTG	TLSSTLGSSL	TVVCDUDDTA	PSDHIVGENE.	SSLRLAARPK
	451	SYTORRHITE	DEMALALMOST	TIISDALETS	SKHSQLSAKL	QLDYRFLLHK
30	501	VI.SKTMDDED	PFVTFITETR	PLAKNEDHYI	FSIQDAFHSL	NLLKAGIDTS
	551	DYEMIMAKIN	RIHAKLWTTH	LISNTESKPT	FPKTACELSL	PFGKKNTVSL
	601	DMEMINAVAC	WDHMNIRWEW	IGNDNVAMTL	ESLHRSKYSL	IKCDRENFIL
•	651	DASKETDÖPP	DSPLSDHRNL	ILGKLFVRPH	PCWNYRLSLR	YGWHRQDTPN
	021	ITEIQMILGT.	KIFEHWQLYG	VYERREADSR	FFFFLKLDKP	KKPPF*

35 (43) Predicted Omp (CPn0021)

One example of a predicted Omp protein is set forth as SEQ ID NOs: 49 & 50 in WO 02/02606. {GenBank accession numbe gi|4376273|gb|AAD18174.1: 'CPn0021'; SEQ ID NO: 43 below}. Preferred Omp proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) 40 to SEQ ID NO: 43; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 43, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of 45 SEQ ID NO: 43. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 43. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 43. Other fragments omit one or more domains of the protein (e.g. omission of a signal 50 peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

	1	MGLFHLTLFG	LLLCSLPISL	VAKFPESVGH	KILYISTQST	COALATYLEA
_	51	LDAYGDHDFF	VLRKIGEDYL	KQSIHSSDPQ	TRKSTIIGAG	LAGSSEALDV
5	101	LSQAMETADP	LQQLLVLSAV	SGHLGKTSDD	LLFKALASPY	PVIRLEAAVR
	151	LANLKNTKVI	DHLHSFIHKL	PEEIQCLSAA	IFLRLETEES	DAYTRDIJ.AA
	201	KKSAIRSATA	LQIGEYQQKR	FLPTLRNLLT	SASPQDQEAT	LYALGKTKDG
	251	QSYYNIKKQL	QKPDVDVTLA	AAQALIALGK	EEDALPVIKK	OALEERPRAT.
40	301	YALRHLPSEI	GIPIALPIFL.	KTKNSEAKLN	VALALLELGC	DTPKLLEVIT
10	351	ERLVQPHYNE	TLALSFSKGR	TLONWKRVNI	IVPQDPQERE	PI.I.STTPGI.E
	401	EQILTFLFRL	PKEAYLPCIY	KLLASOKTOL	ATTAISFLSH	TSHOEALDLL.
•	451	FQAAKLPGEP	ITRAYADLAI	YNLTKDPEKK	RSLHDYAKKL	TOPPLLETOP
	501	ENQRPHPSMP	YLRYOVTPES	RTKLMLDILE	TLATSKSSED	TDIJITOLMTE
	551	GDAKNFPVLA	GLLIKIVE*			TKUUTQUITE
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(44) Oligopeptide Binding Protein Oppa-1 Lipoprotein (CPn0195)

One example of an oligopeptide binding protein is set forth as SEQ ID NOs: 23 and 24 in WO 02/02606. {GenBank accession number gi|4376466|gb|AAD18348.1: 'CPn0195'; SEQ ID NO: 44 below}. Preferred oligopeptide binding proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 44; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 44, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 44. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 44. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 44. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 44

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	1	MRKISVGICI	TILLSLSVVL	Q GCKESSHSS	TSRGELAINI	RDEPRSLDPR
	51	QVRLLSEISL	VKHIYEGLVQ	ENNLSGNIEP	ALAEDYSLSS	DGLTYTEKLK
	101	SAFWSNGDPL	TAEDFIESWK	QVATQEVSGI	YAFALNPIKN	VRKTOEGHLS
40	151	IDHFGVHSPN	ESTLVVTLES	PTSHFLKLLA	LPVFFPVHKS	ORTLOSKSI.P
40	201	IASGAFYPKN	IKQKQWIKLS	KNPHYYNOSO	VETKTITIHF	TPDANTAAKT.
	251	FNQGKLNWQG	PPWGERIPQE	TLSNLOSKGH	LHSFDVAGTS	WITENTHER
	301	LNNMKLREAL	ASALDKEALV	STIFLGRAKT	ADHLLPTNIH	SYPEHOKOEM
	351	AQRQAYAKKL	FKEALEELQI	TAKDLEHLNL	IFPVSSSASS	LIVOLTREOW
AE .	401	KESLGFAIPI	VGKEFALLQA	DLSSGNFSLA	TGGWFADFAD	PMAFI.TTFAV
45	451	PSGVPPYAIN	HKDFLEILQN	IEOEODHOKR	SELVSOASLY	LETERTTEDT
	501	YHDAFQFAMN	KKLSNLGVSP	TGVVDFRYAK	EN*	
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(45) CHLPS 43 kDa Protein Homologue-1 (CPn0562)

One example of a CHLPS protein is set forth as SEQ ID NO: 45 below. GenBank Accession No. GI:4376854; AAD18702.1. Preferred CHLPS proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 45; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 45, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more).

These CHLPS proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 45. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 45. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 45. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

10 SEQ ID No 45

15	121 181 241 301	RVCLQYDEVF ILIFNYPGVM SKEIADGSDS	VLQKICQNFI IDGLELRLPN KSQGNITRNN VRWFVVKDRG NLIGDGLFKK	LLGAGGWIFR AKPDRWMLIS VVKSYQACVR ARSTGAVAKO	PICRDSNLLR NGNSDCLEYR YLRDEPAGPQ FIGSLGWIA	QAYAARLFSA TVLQGEKDWI ARQIVAYGYS	WGVVKFLLGL SFQDHVSSVR FRIAEESQSN LGASVQAEAL KRSKDLHCPE HDHILSDDVI
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(46) YscJ (Yop translocation J protein) (CPn0828)

One example of a YscJ protein is set forth as SEQ ID NO⁵: 109 and 110 in WO 02/02606. {GenBank accession number gi|4377140|gb|AAD18965.1| 'CPn0828'; SEQ ID NO: 46 below. Preferred YscJ proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 46; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 46, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These YscJ proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 46. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 46. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 46. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 46

40	1	MVRRSISFCL	FFLMTLLCCT	SCNSRSLIVH	GLPGREANET	VVLLVSKGVA
•	51	AQKLPQAAAA	TAGAATEQMW	DIAVPSAOIT	EALAILNOAG	LPRMKGTSLI.
	101	DLFAKQGLVP	SELQEKIRYQ	EGLSEOMAST	IRKMDGVVDA	SVOISETTEN
	151	EDNLPLTASV	YIKHRGVLDN	PNSIMVSKIK	RLIASAVPGL	VPENVSVVSD
45	201	RAAYSDITIN	GPWGLTEEID	YVSVWGIILA	KSSLTKFRLT	FYVI.TI.TI.EV
40	251	ISCGLLWVIW	KTHTLIMTMG	GTKGFFNPTP	YTKNALEAKK	AEGAAADKEK
	301	KEDADSQGES	KNAETSDKDS	SDKDAPEGSN	EIEGA*	

(47) Hypothetical (CPn 0415)

One example of a hypothetical protein is set forth as SEQ ID NO^s: 101 and 102 in WO 02/02606. {GenBank accession number gi|4376696|gb|AAD18559.1| 'CPn0415'; SEQ ID NO: 47 below}. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 47; and/or (b) which is a fragment of at

least n consecutive amino acids of SEQ ID NO: 47, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 47. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 47. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 47. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 47

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15	1	MTLIPVIIIV	WCNAFLIKLC	VIMGLOSRLO	HCIEVSONSN	FDSOURORTY
10	51	ACQUETLEQS	VLKIFRYHPL	LKIHDIARAV	YI.I.MAT.EEGE	DIGI.CELMIO
	101	QYPSGAVELF	SCGGFPWKGL	PYPAEHAEFG	LILLOTARRY	PPCONVICTOR
	151	SHFQQALFDH	QGSVFPSLWS	OENSRLLKEK	TTLSOSET.FO	I.CMOTUDEVO
	201	TEDPATGEMM	QRTRSSSAFV	`AASGCOSSLG	AYSSGDVGVT	AVGDCCCDIC
20	251	DCXXEGCCGI	AKEFVCQKSH	OTTEISFLTS	TGKPHPRNTG	FSVI.PDevini
20	301	PLIKCKLLIS	DKQYRVHAAL	AEATSAMTFS	IFCKGKNCOV	VDCDDI.DCCC
	351	LDSYKGPGND	IMILGENDAI	NIVSASPYME	IFALOGKEKF	WNADFLINTE
	401	YKEEGVMLIF	EKKVTSEKGR	FFTKMN*		

(48) Hypothetical (CPn0514)

25 One example of a hypothetical protein is set forth as SEQ ID NOs: 87 and 88 in WO 02/02606. {GenBank accession number gi|4376802|gb|AAD18654.1| SEQ ID NO: 48 below}. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 48; and/or (b) which is a fragment of at least n30 consecutive amino acids of SEQ ID NO: 48, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 48. Preferred fragments of (b) comprise an 35 epitope from SEQ ID NO: 48. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 48. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a 40 transmembrane domain, or of an extracellular domain).

SEQ ID No 48

SYINSF PLSLQLIKRN DIRCVLAPPA DLLNLLIEGK NLGYVP GFGIAANQRI LSVNLYAAPT FFNSPQPRIA LCRHLW RIPTPHILRF ITTKVLRQTP ENYDGLLLIG YDLASG WYDLTKLPFV FALLLHSTSW KEHPLPNLAM LKEAHQ HTGLPPSLLQ EYYALCQYRL GEEHYESFEK
֜֝֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜

(49) Hypothetical (CPn0668)

One example of a hypothetical protein is set forth as SEQ ID NOs: 57 and 58 in WO 02/02606. {GenBank accession number gi|4376968|gb|AAD18807.1 'CPn0668'; SEQ ID NO: 49 below). Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 5 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 49; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 49, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These 10 hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 49. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 49. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the 15 N-terminus of SEQ ID NO: 49. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 49

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1	MKFLLYVPLL	LVLVSTGCDA	KPVSFEPFSG	KLSTOREEDO	HENERVECOO
51	QEFLKKGNFR	KALLCFGIIT	HHEPROTTEN	ONOVITOUCY	MOVED TEST
101	KAFASYLOLP	DAEYSEELFO	MKVATAODEA	CAGINIGACI	FIQUAPULAD
L 51	EDALRIYDEI	LTAPPSKDLG	MAINUMENT AND LIGHT	CGVKKKKICKT	EGFPKLMNAD
201	FPLHILSSEA	FVRISETVIO	VÄVITSIVATI	DIVINDUTEA	TKTLKKLTLQ
251	LNEVVSANVG	AMPEHVADOL.	ZHIVE PHINDQ	IDHFAKLNEE	AMKKQHPNHP
01	LI-VAKCOKRI.	DDICKRAC+	INIGREIERK	KKAEAANIYY	RTAITNYPDT
	101 151 201	101 KAFASYLQLP L51 EDALRIYDEI 201 FPLHILSSEA 251 LNEVVSANVG	CEPLREGNER KALLCEGIIT LO1 KAFASYLQLP DAEYSEELFQ LS1 EDALRIYDEI LTAFPSKDLG CO1 FPLHILSSEA FVRLSEIYLQ LNEVVSANVG AMREHYARGL	CEPLRKGNFR KALLCFGIIT HHFPRDILRN LOI KAFASYLQLP DAEYSEELFQ MKYAIAQRFA LOI EDALRIYDEI LTAFPSKDLG AQALYSKAAL COI FPLHILSSEA FVRLSEIYLQ QAKKEPHNLQ LNEVVSANVG AMREHYARGL YATGRFYEKK	QEFLKKGNFR KALLCFGIIT HHFPRDILRN QAQYLIGVCY KAFASYLQLP DAEYSEELFQ MKYAIAQRFA QGKRKRICRL EDALRIYDEI LTAFPSKDLG AQALYSKAAL LIVKNDLTEA FPLHILSSEA FVRLSEIYLQ QAKKEPHNLQ YLHFAKLNEE LNEVVSANVG AMREHYARGL YATGRFYEKK KKAEAANIYY

(50) Hypothetical (CPn0791)

One example of a hypothetical protein is set forth as SEQ ID NOs: 123 and 124 in 30 WO 02/02606. {GenBank accession number gi|4377101|gb|AAD18929.1| 'CPn0791'; SEQ ID NO: 50 below}. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 50; and/or (b) which is a fragment of at least n35 consecutive amino acids of SEQ ID NO: 50, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 50. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 50. Other preferred fragments lack one or more amino 40 acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 50. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). 45

SEQ ID No 50

50	101 151	YPLGPHRHNE HTLALNPQTI	HNYLLHPMSR PLAAKNLNIA AQDREHLLKM LSTIHVRQAA SSGKLSRIVN	RKVANYILAD LKALKENPKL LTALFTYLRO	NGEIDTVKLV KESIKTLFVP DVGSCFATAP	EAIHHLSQC: SYSTIQNLII
	201	KEPKDPNDPI	SSGKLSRIVN	ORETAUPTNI	SCCTGELEVO	LPTINI VND

	251	LVKLSSSPGL	KKAFSAANLI	ETLGDSEAQI	QQLLSHQYLM	OKLONVHETI.
	301	TANDIIKSTL	LHYYQLQEST	VRAIFFKEGL	FSKEQVAFST	OHPRELSETO
	351	RVYHYLHAYE	EAKSAFIHDT	QNPLLKAWEY	TLATLADASO	PTISNHIRLA
-	401	LGWKSEDPHS	LVSLVTHFVE	EEVENIRILV	QOCEOTYHEA	RSOLEYTEGR
5	451	MRNPLNNQDS	QILTMDHMRF	RQELNKALYE	WDSAQEKAKK	FLHLPEFLLS
	501	FYTKQIPLYF	RSSYDAFIQE	FAHLYANAPA	GFRILFTHGR	THPNTWSPTV
	551	SINEFIRFLS	EFFTSTESEL	LGKHAVINLE	KETSRLVHNI	TAMLHTDVFO
	601	EALLTRILEA	YQLPVPPSIL	NHLDQLSQTP	WVYVSGGTVD	TLLLDYFESS
10	651	EPLTLTEKHP	ENPHELAAFY	ADALKDLPTG	IKSYLEEGSH	SLLSSSPTHV
10	701	FSIIAGSPLF	REAWDNDWYS	YTWLRDVWVK	OHODFLODTI	LPOISTVART
	751	ENFCNKYALQ	HVVHDFHDFC	SDHSLTLPEL	YDKGSRFLSS	LFTKDKTVAL.
	801	IYIRRLLYLM	VREVPYVSEQ	QLPEVLDNVS	SYLGISSRIT	YEKERSLIEE
	851	TIPKMTLLSS	ADLRHIYKGL	LMOSYOKIYT	EEDTYLRLTT	AMRHHMIT.AVD
45	901	APLLFADSNW	PSIYFGFILM	PGTTEIDLWK	FNYAGLOGOP	LDNIOELFAT
15	951	SRPWTLYANP	IDYGMPPPPG	YRSRLPKEFF	*	

(51) Hypothetical (CPn0792)

One example of a hypothetical protein is set forth as SEQ ID NOs: 61 and 62 in WO 02/02606. {GenBank accession number gi|4377102|gb|AAD18930.1| 'CPn0792'; SEQ ID NO: 51 below}. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 51; and/or (b) which is a fragment of at least nconsecutive amino acids of SEQ ID NO: 51, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 51. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 51. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 51. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 51

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		1	MKHTFTKRVL	FFFFLVIPIP	LLLNLMVVGF	FSFSAAKANL	VOVIHTRATN
		51	LSIEFEKKLT	IHKLFLDRLA	NTLALKSYAS	PSAEPYAOAY	NEMMAT.SNTD
40	:	101	FSLCLIDPFD	GSVRTKNPGD	PFIRYLKOHP	EMKKKLSAAV	GKARLLTIDG
40		151	KPLLHYLILV	EDVASWDSTT	TSGLLVSFYP	MSFLOKDLEO	SLHITKONTO
		201	LVNKYGEVLF	CAQDSESSFV	FSLDLPNI.PO	FOARSDSATE	TEKNEGILCO
		251	ENLITVSINK	KRYLGLVLNK	IPIOGTYTLS	LVDVSDLTOS	VI KADI MACE
	:	301	FYVLAFLLMW	WIFSKINTKI.	NKBI OET TEC	MEY PROCEEDING	MUKANIMICE
	3	351	EFNELGNIFN	CTLLLLINST	EKADIDYHSG	EKLOKELCIL	CCIOCATION
45	4	401	DFPTFPKVTF	SSOHLERROL	SCHENGWTVO	DCCDTLCTT	SSLQSALLSP
	4	451	YLYALSARSL	FLAVASSDVS	LOKICKDAND	Decolor	GLAGDIGLES
•		501	VENDETELL	CI CECS DENO	DOVIDEDIAD	SISKITEGNE	AVVAMTFIKY
	_		VEKDRSLELL	SUSEGAPTME	LQRGESFVRL	PLETHQALQP	GDRLICLTGG
	5	551	EDILKYFSQL	PIEELLKDPL	NPLNTENLID	SLTMMLNNET	EHSADGTLTT
	ϵ	501	LSFS*				
50							

(52) Hypothetical (CPn0820)

One example of a hypothetical protein is set forth as SEQ ID NO^s: 113 and 114 in WO 02/02606. {GenBank accession number gi|4377132|gb|AAD18958.1| 'CPn0820'; SEQ ID NO: 52 below}. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 52; and/or (b) which is a fragment of at

least n consecutive amino acids of SEQ ID NO: 52, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 52. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 52. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 52. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain; or of an extracellular domain).

SEQ ID No 52

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16	1	MCNSIAMKKQ	KRGFVLMELL	MSFTLIALLL	GTLGFWYRKI	YTVOKOKERT
15	51	YNFYIEESRA	YKQLRTLFSM	SLSSSYEEPG	SLFSLIFDRG	VYRDPKTACA
	101	VRASLHHDTK	DQRLELRICN	IKDOSYFETO	RLLSHVTHVV	LSFORNPOPE
	151	KLPETIALTI	TREPKAYPPR	TLTYOFAVCK	t	-01 Q.C.L.D.L.D

(53) Hypothetical (CPn0126)

One example of a hypothetical protein is set forth as SEQ ID NO: 53 below. 20 GenBank Accession No. GI:4376390; AAD18279.1 Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 53; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 53, wherein n is 7 or 25 more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 53. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 53. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-30 terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 53. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 53

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40 PDLLDLEDAS ERLRVKASRS LASLPKEISQ LESYIRSAAN DLNTIKTWPH KDQRLVETVS 121 RKLERLAAAQ NYMISELCEI SEILEEEEHH LILAQESLEW IGKSLFSTFL DMESFLNLSH 181 LSEVRPYLAV NDPRLLEITE SEWEVVSHFI NVTSAFKKAQ ILFKNNEHSR MKKKLESVQE 241 LLETFIYKSL KRSYRELGCL SEKMRIHDN PLFPWVQDQQ KYAHAKNEFG EIARCLEEFE 301 KTFFWLDEEC AISYMDCWDF LNESIQNKKS RVDRDYISTK KIALKDRART YAKVLLEENP 361 TTEGKIDLQD AQRAFERQSQ EFYTLEHTET KVRLEALQQC FSDLREATNV RQVRFTNSEN 421 ANDLKESFEK IDKERVRYQK ERLSTVER KNLRDVEAHL EDATMDFEHE VSKSELCSVR ARLEVLEEEL MDMSPKVADI 541 EELLSYEERC ILPIRENLER QKFAIFEKHI QUYSYYEDN GAVVNRKLIN MTERYQNFKR SLNSIQFNGD VLLRDPVYQP 50 721 EGHETRLKER ELQETTLSCK KLKVAODRLS ELESRISRR								
40 121 RKLERLAAAQ NYMISELCEI SEILEEEEHH LILAQESLEW IGKSLFSTFL DMESFLNLSH 181 LSEVRPYLAV NDPRLLEITE SEILEEEHH NVTSAFKKAQ ILFKNNEHSR MKKKLESVQE 241 LLETFIYKSL KRSYRELGCL SEKMRIHDN PLFPWVQDQQ KYAHAKNEFG EIARCLEEFE 301 KTFFWLDEEC AISYMDCWDF LNESIQNKKS RVDRDYISTK KIALKDRART YAKVLLEENP 45 421 ANDLKESFEK IDKERVRYQK EFYTLEHTET KVRLEALQQC FSDLREATNV RQVRFTNSEN 481 RLKGLLRQWK KNLRDVEAHL EQRLYWETID RNEQELREEI GESLRLQNRR KGYRAGYDAG 481 RLKGLLRQWK KNLRDVEAHL EDATMDFEHE VSKSELCSVR ARLEVLEEEL MDMSPKVADI 541 EELLSYEERC ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGUG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERVONFKR SINGLOFMOR WILLERDWINGON		1	MVFSYYCMGL	FFFSGAISSC	${\tt GLLVSLGVGL}$	GLSVLGVLLL	LLAGLLLFKI	OSMIREVPKA
45 121 122 131 132 132 133 134 135 135 136 137 136 137 137 138 138 138 138 138 138	40	9.T	PULLULEDAS	ERLRVKASRS	LASLPKEISO	LESYTRSAAN	DIATT KTWOU	KDODI Mamuro
45 LETFIYKSL KRSYREGCL SEKMRIIHDN PLFPWVQDQQ KYAHAKNEFG EIARCLEEFE 301 KTFFWLDEEC AISYMDCWDF LNESIQNKKS RVDRDYISTK KIALKDRART YAKVLLEENP 45 ADDLKESFEK IDKERVRYQK EFYTLEHTET KVRLEALQQC FSDLREATNV RQVRFTNSEN 481 RLKGLLRQWK KNLRDVEAHL EDATMDFEHE VSKSELCSVR ARLEVLEEEL MDMSPKVADI 541 EELLSYEERC ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGUG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERVONFKR SINGLOFMOD WILLDRINKON.	40	121	KKLEKLAAAQ	NYMISELCEI	SEILEEERHH	T.TT.AOEST.EW	TOVOL DODDE	DMDODT M. C.
45 TIEGKIDLQD AQRAFERQSQ EFYTLEHTET KVRLEALQQC FSDLREATNV RQVRFTNSEN ANDLKESFEK IDKERVRYQK EQRLYWETID RNEQELREEI GESLRLQNRR KGYRAGYDAG ARICKLESPE ANDLKESFEK ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGUG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVNRRLLN MTERVONFKR SINGLOFNCD WILLDRINKON		TOT	LSEVRPYLAV	NDPRLLEITE	ESWEVVSHFI	NVTSAFKKAO	TI.EKAMEUCD	MUZUIT DOMOD
45 TIEGKIDLQD AQRAFERQSQ EFYTLEHTET KVRLEALQQC FSDLREATNV RQVRFTNSEN 421 ANDLKESFEK IDKERVRYQK EQRLYWETID RNEQELREEI GESLRLQNRR KGYRAGYDAG 481 RLKGLLRQWK KNLRDVEAHL EDATMDFEHE VSKSELCSVR ARLEVLEEEL MDMSPKVADI 541 EELLSYEERC ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGVG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERYONFKR SINGLOFMOD WILDERWOOD		241	THETETAYER	KKSYKELGCL	SEKMRIIHDN	PLFPWVODOO	KVAUAKMEEC	ETADOL DEDE
45 421 ANDLKESFEK IDKERVRYQK EFYTLEHTET KVRLEALQQC FSDLREATNV RQVRFTNSEN 421 ANDLKESFEK IDKERVRYQK EQRLYWETID RNEQELREEI GESLRLQNRR KGYRAGYDAG 481 RLKGLLRQWK KNLRDVEAHL EDATMDFEHE VSKSELCSVR ARLEVLEEEL MDMSPKVADI 541 EELLSYEERC ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 661 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGVG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERYONFKP SINGLOFMOD WILDENWOOD		2 O T	KILLMUDEEC	AISYMDCWDF	LNESIONKKS	RVDRDYTSTK	KTAT.KDDADT	VAPULT DESIGN
421 ANDLKESFEK IDKERVRYQK EQRLYWETID RNEQELREEI GESLRLQNRR KGYRAGYDAG 481 RLKGLLRQWK KNLRDVEAHL EDATMDFEHE VSKSELCSVR ARLEVLEEEL MDMSPKVADI 541 EELLSYEERC ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGUG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERYONFKP SLNSTOENGD WLIDDWINGO		2 O T	TIEGKIDLQD	AQRAFEROSO	EFYTLEHTET	KVRLEALOOC	FCDI.DED TOTAL	DOMD DOMOUNT
541 EELLSYEERC ILPIRENLER AYLQYNKCSE ILSKAKFFFP EDEQLLVSEA NLREVGAQLK 601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGVG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERYONEKR SLNSTOENGD MLRDRINGO.	45	421	ANDLKESFEK	IDKERVRYQK	EORLYWETID	RNEOEUREET	GEST.DI.ONDD	VCVD A CVDA C
601 QVQGKCQERA QKFAIFEKHI QEQKSLIKEQ VRSFDLAGVG FLKSELLSIA CNLYIKAVVK 661 ESIPVDVPCM QLYYSYYEDN EAVVRNELLN MTERYONEKE SLNSTOENGD VLIBRANKON		481	RLKGLLRQWK	KNLRDVEAHL	EDATMOFEHE	VSKSELCSVP	ADI.EVI.EPET	MOMODIAINO
661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERYONEKR SLNSTOFNED VILLERRYNDE.		541	EELLSYEERC	ILPIRENLER	AYLOYNKOSE	TICKAKEEED	EDECT TREES	MUNSPRVADI
661 ESIPVDVPCM QLYYSYYEDN EAVVRNRLLN MTERYONFKR SINGIOFNED WILDDRIVED.		601	QVQGKCQERA	OKFAIFEKHI	OEOKSLIKEO	VPSEDIACVC	ELVORITOTA	MUREVGAQUE
721 EGHETRLKER ELOETTLSCK KLKVAODRIS ELESPISER		661	ESIPVDVPCM	OLYYSYYEDN	EAVVENDILLM	MTEDVONEVE	LUKSEUUSTA	CNLYIKAVVK
	50	721	EGHETRLKER	ELOETTLSCK	KIKVAODRIS	FI.FCDI.CDD	STWSTOLMED	APPKDBAAÕB

(54) Hypothetical (CPn0794)

One example of a hypothetical protein is set forth as SEQ ID NO: 54 below. GenBank Accession No. GI:4377105; AAD18932.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more

identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 54; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 54, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 54. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 54. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 54. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 54

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1 MSLYQKWWNS QLKKSLCYST VAALIFMIPS QESFADSLID LNLGLDPSVE CLSGDGAFSV 61 GYFTKAGSTP VEYQPFKYDV SKKTFTILSV ETANQSGYAY GISYDGTITV GTCSLGAGKY 121 NGAKWSADGT LTPLTGITGG TSHTEARAIS KDTQVIEGFS YDASGQPKAV QWASGATTVT 181 QLADISGGSR SSYAYAISDD GTIIVGSMES TITRKTTAVK WVNNVPTYLG TLGGDASTGL 241 YISGDGTVIV GAANTATVTN GNQESHAYMY KDNQMKD

(55) Hypothetical (CPn0796)

One example of a hypothetical protein is set forth as SEQ ID NO: 55 below. GenBank Accession No. GI:4377107; AAD18934.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more 25 identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 55; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 55, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 30 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 55. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 55. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 55. Other fragments omit one or more 35 domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 55

1 MQPCLNMSIV RNSALPLPCL SRSETFKKVR SHMKFMKVLT PWIYRKDLWV TAFLLTAIPG
61 SFAHTLVDIA GEPRHAQAT GVSGDGKIVI GMKVPDDPFA ITVGFQYIDG HLQPLEAVRP
121 QCSVYPNGIT PDGTVIVGTN YAIGMGSVAV KWVNGKVSEL PMLPDTLDSV ASAVSADGRV
181 IGGNRNINLG ASVAVKWEDD VITQLPSLPD AMNACVNGIS SDGSIIVGTM VDVSWRNTAV
241 QWIGDQLSVI GTLGGTTSVA SAISTDGTVI VGGSENADSQ THAYAYKNGV MSDIGTLGGF
301 YSLAHAVSSD GSVIVGVSTN SEHRYHAFQY ADGQMVDLGT LGGPESYAQG VSGDGKVIVG
361 RAQVPSGDWH AFLCPFQAPS PAPVHGGSTV VTSQNPRGMV DINATYSSLK NSQQQLQRLL
421 IQHSAKVESV SSGAPSFTSV KGAISKQSPA VQNDVQKGTF LSYRSQVHGN VQNQQLLTGA
481 FMDWKLASAP KCGFKVALHY GSQDALVERA ALPYTEQGLG SSVLSGFGGQ VQGRYDFNLG
501 TLGVERDLNS HIDEFKGSVS AMGNFVLENS TVSVLRPFAS LAMYYDVRQQ QLVTLSVVMN

(56) Hypothetical (CPn0797)

One example of a hypothetical protein is set forth as SEQ ID NO: 56 below. GenBank Accession No. GI:4377108; AAD18935.1 Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 56; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 56, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, 10 homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 56. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 56. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 56. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of 15 a transmembrane domain, or of an extracellular domain).

SEQ ID No 56

20	ÔΤ	EGWSEAHAVS	GNGSRIVGAS	GAGOGSVTAV	TWESHITKHI.	CTLCCDACCA	TFTDLELLSK EGISKDGEVV
0.5	181 241 301	GVKWEKGKIK GGSVASAEAI	QLKLLPQGLW SANGKVIVGW	MKDLGTLGAT SEANAISEDG STTNNGETHA	YSVARGVSGD TVIVGRGEIS FMHKDETMHD	GSIIVGVSAT RNHIVAVKWN	ARGEDYGWQV KNAVYSLGTL ATGVSADGRA TDAGAERAYL

(76) Oligopeptide Binding Protein Oppa-2 Lipoprotein (CPn0196)

One example of an oligopeptide binding protein is set forth as SEQ ID NOs: 127 and 30 128 in WO 02/02606. {GenBank accession number GI:4376467; AAD18349.1 'CPn0196'; SEQ ID NO: 76 below}. Preferred oligopeptide binding proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 76; and/or (b) which is a fragment of at 35 least n consecutive amino acids of SEQ ID NO: 76, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO 76. Preferred fragments of (b) 40 comprise an epitope from SEQ ID NO: 76. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 76. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of 45 a transmembrane domain, or of an extracellular domain).

SEQ ID No 76

1 mlrffavfis tlwlitsgcs psqsskgifv vnmkemprsl dpgktrliad qtlmrhlyeg
61 lveehsqnge ikpalaesyt isedgtrytf kiknilwsng dpltaqdfvs swkeilkeda
121 ssvylyaflp iknaraifdd tespenlgvr aldkrhleid letpcahflh fltlpiffpv
181 hetlrnysts feempitcga frpvslekgl rlhleknpmy hnksrvklhk iivqfisnan
241 taailfkhkk ldwdgppwge pippeisasl hdddqlfslp gasttwllfn iqkkpwnnak
301 lrkalslaid kdmltkvvyd glaeptdhil hprlypgtyp erkrqneril eaqqlfeeal

- 361 delqmtredl eketltfstf sfsygricqm lreqwkkvlk ftipivgqef ftiqknfleg 421 nysltvnqwt aafidpmsyl mifanpggis pyhlqdshfq tllikitqeh kkhlrnqlii 481 ealdylehch ileplchpnl rialnknikn fnlfvrrtsd frfiekl
- Seventh Antigen Group

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The immunogenicity of other Chlamydia pneumoniae antigens may be improved by combination with two or more Chlamydia pneumoniae antigens from either the first antigen group or the second antigen group or the third antigen group or the fourth, antigen group or the fifth antigen group or the sixth antigen group. Such other Chlamydia pneumoniae antigens include a seventh antigen group consisting one or more hypothetical proteins (ie proteins which, for example, have no known cellular location and/or function. These antigens are referred to herein as the "seventh antigen group". Each of the Chlamydia pneumoniae antigens of the seventh antigen group is described in more detail below.

(57) Hypothetical (CPn0331)

One example of a hypothetical protein is set forth as SEQ ID NO: 57 below. GenBank Accession No. GI:4376609; AAD18480.1. Preferred hypothetical proteins 20 for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 57; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 57, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 25 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 57. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 57. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 57. Other fragments omit one or more 30 domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEO ID No 57

~-	2221						
35	1	MAVSGGGGVQ	PSSDPGKWNP	ALQGÉQAEGP	SPLKESIFSE	TKQASSAAKO	ESLVRSGSTG
	61	MYATESQINK	AKYRKAQDRS	STSPKSKLKG	TFSKMRASVO	GFMSGFGSRA	SRVSAKRASD
	121	SGEGTSLLPT	EMDVALKKGN	RISPEMOGFF	LDASGMGGSS	SDISOLSLEA	LKSSAFSGAR
	181	SLSLSSSESS	SVASFGSFQK	AIEPMSEEKV	NAWTVARLGG	EMVSSLLDPN	VETSSLVERA
40	241	MATGNEGMID	LSDLGQEEVS	TAMTSPRAVE	GKVKVSSSDS	PEANPTGIPN	SNTLERAEKE
	301	AEKQESREQL	SEDQMMLARA	MAGLLTGAAP	OEVLSNSVWS	GPSTVFPPPK	FSGTLPTOPS
	361	GDKSKHKSPG	IEKSTNHTNF	SPLREGTVKS	AEVKSLPHPE	SMYRFPKDST	VSREEDEAVA
	421	Kestafknpe	NSSQNFLPIA	VESVFPKESG	TGGALGSDAV	SSSYHFIAOR	GVSLTADI.DD
	481	ATDDYKEKLE	AHKGPGGPPD	PLIYOYRNVA	VEPPIVLRSP	OPESGSSRIS	AUGRDEY V GA
	541	HDDGGGGNSG	GFSGDQRRGS	SGOKASROEK	KGKKLSTDI .	R	A GUITTING
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(58) Hypothetical (CPn0234)

One example of a hypothetical protein is set forth as SEQ ID NO: 58 below. GenBank Accession No. gi|4376508|gb| AAD18387.1 Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 5 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 58; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 21, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 58. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 58. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 58. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 58

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- 1 MLQSCKKALL SIVVSILAFH PIPGMGVEAK SGFLGKVKGW FSKKEIQEEA RILPVKDSLS 61 WKRYDYTSSS GFSVEFPGEP DHSGQIVEVP QSEITIRYDT YVTETHPDNT VYVVSVWEYP 121 EKVDISRPEL NLQEGFSGMM QALPESQVLF MQARQIQGHK ALEFWIVCED VYFRGMLISV
 - 181 NHTLYQVFMV YKNKNPQALD KEYEAFSQSF KITKIREPRT IPSSVKKKVS L

(59) Hypothetical (CPn0572)

One example of a hypothetical protein is set forth as SEQ ID NO: 59 below. Genbank 25 Accession No. gi|4376866|gb|; AAD18712.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 59; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 59, wherein n is 7 or more (e.g. 8, 10, 30 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 59. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 59. Other preferred fragments lack one or 35 more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 59. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). 40

SEQ ID No 59

45	OT WOODGOW	PST TTQITQTGQT VSF SAEHSFSTLI	PETGSVGATA	OSBOSBET.TO	TECHTOPPO	Trocass-
40	TET TOVIOON	HOO GETSKYESSE	DLGDLDSLSC	SERAFGARGE	PODOCI DROW	TRITTEDDO
	TOT DITMETIV	MPA VQQKMOTKGC	HEVYVDEARS	SETEMONOM	CUNTCATON	3 7/07/200
	AAT ADDRIVE	HAR CVGYETIHSE) WTGRVKPTME	ERSCATCMVM	UT.MT.CMPTPO	AIRMONING
50	301. DGGGGT	PSA WKKGAKVETG	PIWDDVGGTK	GINWKTTOAD	DECETATIONS	CONTRACTOR
	JOT LGIEAGW	raa bmanaanreet	KVDLGGINLG	CTTTVIVITE	CCCONTENOUR	CMOMPATA
50	421 1010000	TTE EDITORDDEG	OGEDDNATEG	ממשמשתואיי	DMI CCCDT I M	T 0373 65
	TOT ONAKONDI	VIA IDSNGNSVSD	LNODLGOVAK	NSENCUMPOT	WIT DEFINATION	DDOGGATER
	O THE TERROGETI	CAT TOWNTOSTGO	SEGATPTPOD	TTAKTUTCI.D	VANUECCCUT	DODOTTO TOTAL
55	OUT POWKING	IST ISTETETEST	STTSTGTGTG	SVSTOSTOVC		CO A COMPONE OF A
	OOT STOTEOW	LP SGIRHVATIS	IVRNAACRCT	TIT.OOCCDCOC	FPIPPSGTGT	ONMGAOT WAA
99	/21 ASQVASTI	GQ VVNQAATAGS	OPSSRRSSPT	SPRRK		~

Eight Antigen Group

The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group of the third antigen group or the fourth antigen group or the fifth or the sixth antigen group or the seventh antigen group. Such other *Chlamydia pneumoniae* antigens include an eight antigen group consisting one or more FACS positive CPn antigens. These antigens are referred to herein as the "eight antigen group". Each of the *Chlamydia pneumoniae* antigens of the eight antigen group is described in more detail below.

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(60) Low Calcium Response Protein H (CPn0811)

One example of a Low Calcium Response Protein H is set forth as SEQ ID NO: 60 below. Genbank Accession No. GI:4377123; AAD18949.1. Preferred low calcium response proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 60; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 60, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These low calcium response proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 60. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 60. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 60. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 60

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1 mskpsprnan qpqkpsasfn kktrsrlael aaqkkakadd leqvhpvpte eeikkalgni 61 feglsngldl qqilglsdyl leeiytvayt fysqgkynea vglfqllaaa qpqnykymlg 121 lsscyhqlhl yneaafgffl afdaqpdnpi ppyyiadsll klqqpeesnn fldvtmdicg 181 nnpefkilke rcqimkqsie kqmagetkka ptkkpagksk tttnkksgkk r

35 (61) Yop Proteins Translocation Protein T (CPn0823)

One example of a Yop Proteins Translocation Protein T is set forth as SEQ ID NO: 61 below. Genbank Accession No. GI:4377135; AAD18960.1. Preferred Yop proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 61; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 61, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more): These Yop proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 61. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 61. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 61. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

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1 mgislpelfs nlgsayldyi fqhppayvws vfllllarll pifavapflg aklfpspiki 61 gislswlaii fpkvladtqi tnymdnnlfy vllvkemiig ivigfvlafp fyaaqsagsf 121 itnqqgiqgl egatslisie qtsphgilyh yfvtiifwlv gghrivisll lqtlevipih 181 sffpaemmsl sapiwitmik mcqlclvmti qlsapaalam lmsdlflgii nrmapqvqvi yllsalkafm gllfltlaww fiikqidyft lawfkevpim llgsnpqvl

(62) Yop Proteins Translocation Protein J

10 One example of a Yop Proteins Translocation Protein J is set forth as SEQ ID NO: 62 below Genbank Accession No. GI:4377140; AAD18965.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 62; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 62, wherein n is 7 or 15 more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 62. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 62. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-20 terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 62. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). 25

SEQ ID No 62

1 mvrrsisfcl fflmtllcct scnsrslivh glpgreanei vvllvskgva aqklpqaaaa
61 tagaateqmw diavpsaqit ealailnqag lprmkgtsll dlfakqglvp selqekiryq
121 eglseqmast irkmdgvvda svqisftten ednlpltasv yikhrgvldn pnsimvskik
181 rliasavpgl vpenvsvvsd raaysditin gpwglteeid yvsvwgiila kssltkfrli
241 fyvlililfv iscgllwviw kthtlimtmg gtkgffnptp ytknaleakk aegaaadkek
301 kedadsqges knaetsdkds sdkdapegsn eiega

35 (63) OmpA (CPn0695)

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One example of an OmPA encoded (MOMP) protein is set forth as SEQ ID NO: 63 below Genbank Accession No. GI:4376998; AAD18834.1. Preferred OmpA proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 63; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 63, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmpA proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 63. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 63. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 63. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

5 mkkllksall saafagsvgs lqalpvgnps dpsllidgti wegaagdpcd pcatwcdais 61 lragfygdyv fdrilkvdap ktfsmgakpt gsaaanytta vdrpnpaynk hlhdaewftn 121 agfialniwd rfdvfctlga sngyirgnst afnlvglfgv kgttvnanel pnvslsngvv 181 elytdtsfsw svgargalwe cgcatlgaef qyaqskpkve elnvicnvsq fsvnkpkgyk 241 gvafplptda gvatatgtks atinyhewqv gaslsyrlns lvpyigvqws ratfdadnir 301 iaqpklptav lnltawnpsl lgnatalstt dsfsdfmqiv scqinkfksr kacgvtvgat

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(64) Hypothetical (CPn0210)

One example of a Hypothetical Protein is set forth as SEQ ID NO: 64 below Genbank Accession No. GI:4376482; AAD18363.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 64; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 64, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 64. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 64. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 64. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 64

1 mlvelealkr efahlkdqkp tsdqeitsly qcldhlefvl lglgqdkflk atededvlfe
61 sqkaidawna lltkardvlg lgdigaiyqt ieflgaylsk vnrrafcias eihflktair
121 dlnayylldf rwplckieef vdwgndcvei akrklctfek etkelnesll reehamekcs
181 iqdlqrklsd iiielhdvsl fcfsktpsqe eyqkdclyqs rlryllllye ytllcktstd
241 fqeqarakee firekfslle lekgikqtke lefaiakskl ergclvmrky eaaakhslds
301 mfeeetvksp rkdte

(65) Low Calcium Response Locus Protein H (CPn1021)

One example of a Low Calcium Response Protein H is set forth as SEQ ID NO: 65 40 below Genbank Accession No. GI:4377352; AAD19158.1. Preferred low calcium response proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 65; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 65, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 45 100, 150, 200, 250 or more). These low calcium response proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 65. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 65 Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 50 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 65. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

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1 mshlnyllek iaasskedfp fpddlesyle gyvpdknial dtyqkifkis sedlekvyke 61 gyhayldkdy aksitvfrwl vffnpfvskf wfslgaslhm seqysqalha ygvtavlrdk 121 dpyphyyayi cytltnehee aekalemawv raqhkplyne lkeeildirk hk

Ninth Antigen Group

The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group or the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group. Such other *Chlamydia pneumoniae* antigens include a ninth antigen group. These antigens are referred to herein as the "ninth antigen group". Each of the *Chlamydia pneumoniae* antigens of the ninth antigen group is described in more detail below.

(66) Low Calcium Response Protein D (CPn0323)

One example of a Low Calcium Response Protein D is set forth as SEQ ID NO: 66 below Genbank Accession No. GI:4376601; AAD18472.1. Preferred low calcium response proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 66; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 66, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These low calcium response proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 66. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 66. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 66. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 66

1 mnkllnfvsr tlggdtalnm inkssdlila lwmmgvvlmi iiplpppivd lmitinlsis
61 vfllmvalyi psalqlsvfp slllittmfr lginisssrq illkayaghv iqafgdfvvg
121 gnyvvgfiif liitiiqfiv vtkgaervae vaarfrldam pgkqmaidad lragmidatq
181 ardkraqiqk eselygamdg amkfikgdvi agivislini vggltigvam hgmdlaqaah
181 vytllsigdg lvsqipslli altagivttr vssdkntnlg keistqlvke ggsgasttvg agggasttvg agggasttvg agggasttvg ggsgasttvg ggsgasttvg skliqhktks gqsfvddmip kmrqalyqdi girypgihvr vrgkipphhv ltnevednls rynlpfityk naaglpsawv vilhlsyff hkssqeflgi qevrsmiefm ersfpdlvke
181 sedakailek aaikywtple viilhlsyff hkssqeflgi qevrsmiefm ersfpdlvke ggspvlltai dvrryvrkli etefpdiavi syqeilpeir iqplgriqif

(67) CHLPS 43kDa Protein Homolog-1 (CPn0062)

One example of a CHLPS 43kDa Protein Homolog-1 is set forth as SEQ ID NO: 67 below Genbank Accession No. GI:4376318; AAD18215.1. Preferred CHLPS proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 67; and/or (b) which is a

fragment of at least n consecutive amino acids of SEQ ID NO: 67, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These CHLPS proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 67. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 67. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 67. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 67

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15		l mmsskrtski	avlsilltft	hsigfanans	svalatvvit	sevvkkpaka	gerkaskken
10	U.	ratingytyps	sitisaradk	mknssrkess	aacheigang	transcrated assess	Inner 1 .
		dragramit.	KETTDKTDSK	GKtsiherek	aterfimeer] cccamleane	
	TO.	recerviable	rekerdaner	hipvvavatn	nkeantkttk	al samed and	
	24.	radagastbar	vrsnpevsva	rokeellkel	Vaerrockrk	CTTTCTTCTTC	1 41-1
00	30.	. veserryape	Kaaelksrrn	CKVspeared	kvssckrdar	andleadles sa	
20	30.	. cgagivikib	Ksqvasnaon	fvrnskntni	devitance		secasqeeqq
	421	nsisvctmvv	tviamivgal	iianatesqt	tsdptpptpt	caseerawbc	sscvskrrtn

(68) Hypothetical (CPn0169)

One example of a CHLPS 43kDa Protein Homolog-1 is set forth as SEQ ID NO: 68 25 below Genbank Accession No. GI:4376437; AAD18322.1. Preferred CHLPS proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 68; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 68, wherein n is 7 or 30 more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These CHLPS proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 68. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 68. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-35 terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 68. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

40 SEQ ID No 68

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1 mknvgsecsq plvmelntqp lrnlcesrlv kitsfviall alvggitlta lagagilsfl
61 pwlvlgivlv vlcalfllfs ykfcpikelg vvyntdsqih qwfqkqrnkd lekatenpel
121 fgenraednn rsarsqvket lrdcdgnvlk kiyernldvl lfmnwvpktm ddvdpvseds
181 irtviscykl ikackpefrs lisellramq sglgllsrcs ryqeraktvs hkdaplfcpt
241 hsyyrdgylt plragpryii nrai

(69) PmpD family (CPn0963)

One example of a PmpD protein is set forth as SEQ ID NO: 69 below Genbank Accession No. GI:4377287; AAD19099.1. Preferred PmpD proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 69; and/or (b) which is a fragment of at

least *n* consecutive amino acids of SEQ ID NO: 69, wherein *n* is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These PmpD proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 69. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 69. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 69. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 69

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15	121 181 241	klvwsdldtr spetavaais lgfenlkapk	nfsqptqepd edleisenis sgaavysdrd	ytqvpltqgs tsnavsekis ardplqglaf ivfenlykgl	sgesldlada sdtkenrkdl fykntssqsi	lgvfnkqfee nflehfqhlf etedpskksg sekdssfqgi gsaagvnivv vvgnkgaivv	eettvfgidq lkevssdlpk ifsgsgansg
20	421	gltdfvggga	laaqgtltlr	nnavvqcvkn	ssasdidiqg	ncsaiefsgn	qslialgehi
	481	altggalsan	dkviiannfg	eilfeqnevr	tskthggail	agtvdlneti	sevafkqnta
	541	sgaitflknk	asvlevmtga	edvaggalw	nhggaiycgc	rsnpkleqkd	sgeniniign
25	661	shgdhyppkt	veeevppsll	eehpvvsstd	aqkyvapqet	apvesdasst	nkdekslnac
	721	eesstvgdla	ivgggallst	nevnvcsnqn	irgggailaq	hifitdntgn	lrfsgnlggg
	781	vefvsngsgk	fggavcalne	svnitdngsa	vvfsdnvtsn	gcdsggaila	kkvdisanhs
	841	iafkenfvfg	sengrsgga	iianssvnid	vsfsknrtrl	ggagvaapqg	svticgnqgn
30	961 1021 1081 1141	psgagvqiad qdaityeent qegtlalsqn vqinmssptp	ggtvcleafg irglpdkdvs aelwlaglkq nkdkavdtpv	stqtaasise gdilfegnin plsapslifn etgssivlsa ladiisityd	kdsfgggaiy fdgsfnaihl skpqddsaqh gsilrifdsq	tqnlkivkna cgndskivel hegtirfsrg vdssaplpte	gnvsfygnra savqdkniif vskipqiaai nkeetlvsag
35	1261	atyghtgvws	eskmedgrlv	vgwqptgykl	ktaegmtgtp	tadaslsnik	idvslpsitp
	1321	htiagrmeld	fstnvwgsgl	gvvedcqnig	npekggalvl	nnlwshytdl	ralkqeifah
	1381	sqffgktesq	sykakndvks	ymgaayagil	efdgfkhhlt	gyalgldtql	vedfliggcf
	1441	gswigkgfia	gtsidyrviv	nprrfisaiv	agpwlikgaf	vygninndlt	tdygtlgist
40		vdipckawka	rgratenays	rgsraevnsv	alavyfdwyr	kanzali+ll-	daayswksyg

Tenth Antigen Group

The immunogenicity of other *Chlamydia pneumoniae* antigens may be improved by combination with two or more *Chlamydia pneumoniae* antigens from either the first antigen group or the second antigen group or the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group or the ninth antigen group. Such other *Chlamydia pneumoniae* antigens include a tenth antigen group. Each of the *Chlamydia pneumoniae* antigens of the tenth antigen group is described in more detail below.

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(70) OmpH-like outer membrane protein (CPn0301)

One example of 'OmpH-like' protein is disclosed as SEQ ID NOs: 77 & 78 in WO 02/02606. {GenBank accession number: gi|4376577|gb|AAD18450.1| 'CPn0301'; SEQ ID NO: 70 below and SEQ ID No 4 above}. Preferred OmpH-like proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 4; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 3, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmpH-like proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 4. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 4. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 19 or more, to remove the signal peptide) from the N-terminus of SEQ ID NO: 4. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

30 SEQ ID No 70

- 1 MKKLLFSTFL LVLGSTSAAH ANLGYVNLKR CLEESDLGKK ETEELEAMKQ
- 51 QFVKNAEKIE EELTSIYNKL QDEDYMESLS DSASEELRKK FEDLSGEYNA 101 YQSQYYQSIN QSNVKRIQKL IQEVKIAAES VRSKEKLEAI LNEEAVLAIA

35 151 PGTDKTTEII ALLNESFKKQ N*

(71) L7/L12 Ribosomal Protein (CPn0080)

One example of an L7/L12 Ribosomal protein is set forth as SEQ ID No 71 below (GenBank accession number: GI:4376338; AAD18233.1). 'CPn0080'; SEQ ID NO: 71 below. Preferred L7/L12 proteins for use with the invention comprise an 40 amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 71; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 71, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 45 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These L7/L12 ribosomal proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 71. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 71. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 19 or more, to remove 50 the signal peptide) from the N-terminus of SEQ ID NO: 71. Other fragments omit one

or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 71

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1 mttesletlv eklsnltvle lsqlkkllee kwdvtasapv vavaagggge apvaaeptef 61 avtledvpad kkigvlkvvr evtglalkea kemteglpkt vkektsksda edtvkklqda 121gakasfkgl

10 (72) AtoS two-component regulatory system sensor histidine kinase protein (CPn0584)

One example of 'AtoS' protein is disclosed as SEQ ID NOs: 105 & 106 in WO 02/02606. {GenBank accession number: gi|4376878|gb|AAD18723.1| 'CPn0584'; SEQ ID NO: 72 below and SEQ ID No 9 above}. Preferred AtoS proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 72; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 72, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more).

These AtoS proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 72. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 72. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 72. Other fragments omit one or more domains of the

protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEQ ID No 72

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1	MNVPDSKNLH	PPAYELLEIK	ARITOSYKEA	SATITATIONS	TT.T.T.CETCTITE
51	DICNSQAREL	LGIDENLEIL	NRSFTDVLPD	TCLCESTOEN	LECT PURCHE
10:	RLSLCKESKE	KEVELFIRKN	EISGYLFTOT	DUDGUAROTE	DESDEVANTE
151	ELGKMTATLA	HEIRNPLSGI	VGFASTLKKE	THEORIGISST	NAIERIKNIA
201	NNLVSSMLEY	TKSQPLNLKI	INLODEESSI.	TDI.I.CUCROM	SSIISGIRSL
251	LFRSIDPDRM	NSVVWNLVKN	AVETGNSDIT	LTLUTCONTO	CKFVKEGAQP
301	IMDKLFTPFF	TTKREGNGLG	TARACKTTRI.	DIDDIOGOIS	VINPGTIPSE
351	PELLAALPKE	RAAS*	THING WELL	HGGDIQUKIS	DSAVSFFIII

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(73) OmcA 9kDa-cysteine-rich lipoprotein(CPn0558)

One example of 'OmcA' protein is disclosed as SEQ ID NOs: 9 & 10 in WO 02/02606. {GenBank accession number: gi|4376850|gb|AAD18698.1| 'CPn0558', 'OmcA', 'Omp3'; SEQ ID NO: 73 below and SEQ ID No 10 above}. Preferred OmcA proteins for use with the invention comprise an amino acid sequence: (a) 5 having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 73; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 73, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These OmcA proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 73. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 73. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more; preferably 18 or more to remove the signal peptide) from the N-terminus of SEQ ID NO: 73. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide as described above, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain). The protein may be lipidated (e.g. by a N-acyl diglyceride), and may thus have a N-terminal cysteine.

SEQ ID No 73

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MKKAVLIAAM FCGVVSLSSC CRIVDCCFED PCAPSSCNPC EVIRKKERSC GGNACGSYVP SCSNPCGSTE CNSQSPQVKG CTSPDGRCKQ *

(74) Hypothetical (CPn0331)

One example of a hypothetical protein is set forth as SEQ ID NO: 74 below and SEQ ID No 57 above. Genbank Accession No. GI:4376609; AAD18480.1. Preferred hypothetical proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 74; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 74, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO: 74. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 74. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 74. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

SEO ID NO 74

	226171101	**				
45	121 sge	vsggggvq pssdpgkwn atesqink akyrkaqdr: egtsllpt emdvalkkgn slsssess svasfosfo	s sispkskikg 1 rispemaaff	tiskmrasvq	gfmsgfgsra	srvsakrasd
50	301 aek 361 gdk 421 kes 481 atd	gnegmid lsdlgqeevergesreql sedgmmlaraskhkspg iekstnhtnistafknpe nssqnflpiaddykekle ahkgpggppd	maglitgaap spiregtvks vesvfpkesg	gkvkvsssds qevlsnsvws aevkslphpe tggalgsdav	peanptgipn gpstvfpppk smyrfpkdsi	sntleraeke fsgtlptqrs vsreepeavv

(75) PmpD family (CPn0963) ·

One example of a PmpD protein is set forth as SEQ ID NO: 75 below Genbank Accession No. GI:4377287; AAD19099.1. Preferred PmpD proteins for use with the invention comprise an amino acid sequence: (a) having 50% or more identity (e.g. 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more) to SEQ ID NO: 75; and/or (b) which is a fragment of at least n consecutive amino acids of SEQ ID NO: 75, wherein n is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These hypothetical proteins include variants (e.g. allelic variants, homologs, 10 orthologs, paralogs, mutants, etc.) of SEQ ID NO: 75. Preferred fragments of (b) comprise an epitope from SEQ ID NO: 75. Other preferred fragments lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the Cterminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the N-terminus of SEQ ID NO: 75. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of 15 a transmembrane domain, or of an extracellular domain).

SEQ ID No 75 :

20	1	muskkturen	200 E-b				•
	61	mvakkevisy	rssisnsviv	allsagiafe	ahslhsseld	lgvfnkqfee	hsahveeaqt
	121	kluwediden	psqxesexvi	Acdabredas	sgesldlada	nflehfqhlf	eettvfgidq
	181	Enetarrasia	ursqprqepa	tsnavsekis	sdtkenrkdl	etedpskksg	lkevssdlpk
	247	lafenlkank	edieisenis	ardpidglaf	fykntssqsi	sekdssfqgi	ifsgsgansg
25		TATEITIVODY	suaavvsara	10161111111	eficacalod	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4.1
	201	carginiear	LIVKGISTGG	avrtarnhev	mplaceile	7 P7	- 1
	204	Baracapray	Simientalik	engalsogai	- acaedidiaa	naaniafama	
	- T-22 I	great vyggga	raaddcitirr	ทุกลงงงงงงงหา	tekthogasil		
		artggarsan	ukviiannig	elltemevr	nhaasiwaaa	Lalma Inlamor	
30		ハスペットナインコン	asvievillida	egvagggalw	appare I dama		
••	001	agarracur	veisinisday	VIKankaaci	ackizzance	222222222	-1-2-1-3
	001	PridomAbbyr	veeevppsii	eenpvvsstd	iraaaailaa	hifitantan	7 6 7
	12.1	eesscvgara	ivaddatist	nevnycanon	wefednyten	ccdccc-il-	1-11-1
	, 0 -	ACTABITGEOV	rqqavcarne	SVD1Ednosa	vefeknytyl	~~~~~	
35	0.4.7	TOTVETITATO	senarsadaa	11anggimia	dnacdil for		-
.00	201	Subrerefed	usquiitakn	SECTARASISA	kde faranin	to area 7 la d'a alarea a	
	201	pagagvgrad	ddcvcreard	adilteanin	fdacfnaihl	condoled1	
	.021	dogreate	TIGIDORGAS	DISABSITED	cknodddaawh	how-d	
	T00T	dedcrargon	germragiku	Prassivies	acilyifda.		
40	****	AdTHURSPOLD	iikukavatby	ladiisitvd	leefimaada	+1~1~~~ <i>idd</i>	-117 99
	1201	TOTETTOPLII	vovennatis	Shkdiblisi	ktaecmtata	6-d1	2 3
	1201	acygnicgows	eskilledariv	Vawantavki	nnekaal	~~1 h	7 1 1 C - 1
		TYC TOOL METO	Tachiamagai	avvencania	ofdofbhhle	~~.~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	7071
	TOOT	pdrraveerd	Sykaknovks	vmgaavagi i	acowlikesf	75777	والمستحدث والمستحدث والمستحدث
45		aparavarra .	Great GALATA	DDITIISALV	Stimmfiges	raceidle.	
.0	#30T	verremogrb	rgratenays	rgsraevnsv	alayyfdyner	bommen 1 i + 1 l-	daayswksyd
	1201	vdipckawka	risnntewns	ylstylafny	ewredliayd	fnggiriif	

Preferably the composition of the invention comprises a combination of CPn antigens selected from the group consisting of: (1) CPn0301 and CPn0080; (2) CPn 0584 and CPn 0558; and (3) CPn 0331 and CPN 0963. Preferably the composition comprises a combination of any one or more of groups (1), (2) and (3).

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Even more preferably, the composition of the present invention comprises a combination of CPn antigens selected from the group consisting of (1) CPn0385, CPn0324, CPn 0503, CPn0525 and CPn 0482. Preferably the composition is administered in the presence of alum and/or cPG.

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The invention thus includes a composition comprising a combination of *Chlamydia pneumoniae* antigens, said combination selected from the group consisting of two, three, four, five or six *Chlamydia pneumoniae* antigens of the first antigen group and two, three, four, five, or six *Chlamydia pneumoniae* antigens of the second antigen group. Preferably, the combination is selected from the group consisting of three, four, five or six *Chlamydia pneumoniae* antigens from the first antigen group and three, four, five or six *Chlamydia pneumoniae* antigens from the second antigen group. Still more preferably, the combination consists of six *Chlamydia pneumoniae* antigens from the first antigen group and three, four, five or six, *Chlamydia pneumoniae* antigens from the second antigen group.

The invention further includes a composition comprising a combination of Chlamydia pneumoniae antigens, said combination selected from the group consisting of two, three, four, five or six, Chlamydia pneumoniae antigens of the second antigen group and two, three, four, five, six, seven or eight Chlamydia pneumoniae antigens of the third antigen group. Preferably, the combination is selected from the group consisting of three, four, five or six Chlamydia pneumoniae antigens from the second antigen group and three, four, five, six, seven or eight Chlamydia pneumoniae from the third antigen group. Still more preferably, the combination consists of six Chlamydia pneumoniae antigens from the second antigen group and three, four, five, six, seven or eight Chlamydia pneumoniae antigens of the third antigen group.

There is an upper limit to the number of *Chlamydia pneumoniae* antigens which will be in the compositions of the invention. Preferably, the number of *Chlamydia pneumoniae* antigens in a composition of the invention is less than 20, less than 19, less than 18, less than 17, less than 16, less than 15, less than 14, less than 13, less than 12, less than 11, less than 10, less than 9, less than 8, less than 7, less than 6, less than 5, less than 4, or less than 3. Still more preferably, the number of *Chlamydia pneumoniae* antigens in a composition of the invention is less than 6, less than 5, or less than 4. The *Chlamydia pneumoniae* antigens used in the invention are preferably isolated, i.e., separate and discrete, from the whole organism with which the molecule is found in nature or, when the polynucleotide or polypeptide is not found in nature, is sufficiently free of other biological macromolecules so that the polynucleotide or polypeptide can be used for its intended purpose.

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In either of the above combinations, preferably the composition comprises one or more *Chlamydia pneumoniae* antigens from the fourth antigen group which includes porB. Or, alternatively, in either of the above combinations, preferably the *Chlamydia pneumoniae* antigens from the fourth antigen group includes one or more members of the pmp3 family.

Other aspects of the present invention are presented in the accompanying claims and in the following description and drawings. These aspects are presented under separate section headings. However, it is to be understood that the teachings under each section are not necessarily limited to that particular section heading.

Before describing the present invention in detail, it is to be understood that this invention is not limited to particularly exemplified molecules or process parameters as such may, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments of the invention only, and is not intended to be limiting. In addition, the practice of the present invention will employ, unless otherwise indicated, conventional methods of virology, microbiology, molecular biology, recombinant DNA techniques and immunology all of which are within the ordinary skill of the art. Such techniques are explained fully in the literature. See, e.g., Sambrook, et al., Molecular Cloning: A Laboratory Manual (2nd Edition, 1989); DNA Cloning: A Practical Approach, vol. I & II (D. Glover, ed.); Oligonucleotide Synthesis (N. Gait, ed., 1984); A Practical Guide to Molecular Cloning (1984); and Fundamental Virology, 2nd Edition, vol. I & II (B.N. Fields and D.M. Knipe, eds.).

All publications, patents and patent applications cited herein, whether supra or infra, are hereby incorporated by reference in their entirety. It must be noted that, as used in this specification and the appended claims, the singular forms "a", "an" and "the" include plural referents unless the content clearly dictates otherwise. All scientific and technical terms used in this application have meanings commonly used in the art unless otherwise specified. As used in this application, the following words or phrases have the meanings specified.

The term "comprising" means "including" as well as "consisting" e.g. a composition "comprising" X may consist exclusively of X or may include something additional e.g. X + Y.

The term "about" in relation to a numerical value x means, for example, x±10%.

References to a percentage sequence identity between two amino acid sequences means that, when aligned, that percentage of amino acids are the same in comparing the two sequences. This alignment and the percent homology or sequence identity can be determined using software programs known in the art, for example those described in section 7.7.18 of Current Protocols in Molecular Biology (F.M. Ausubel et al., eds., 1987) Supplement 30. A preferred alignment is determined by the Smith-Waterman homology search algorithm using an affine gap search with a gap open penalty of 12 and a gap extension penalty of 2, BLOSUM matrix of 62. The Smith-Waterman homology search algorithm is disclosed in Smith & Waterman (1981) Adv. Appl. Math. 2: 482-489.

IMMUNE RESPONSE

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The mechanism by which the immune system controls disease includes the induction of neutralising antibodies via humoral immunity and the generation of T-cell responses via cellular immunity. As used herein, the term "immune response" against an antigen refers to the development in a host mammalian subject of a humoral and/or a cellular immune response against that antigen.

As used herein, the term "humoral immune response" refers to an immune response mediated by antibody molecules. The antibodies generated by humoral immunity are primarily effective against extracellular infectious agents.

- 5 SEQ ID Nos 1-76 in the compositions of the invention may be supplemented or substituted with an antibody that binds to the protein. This antibody may be monoclonal or polyclonal.
- As used herein, the term "cell mediated immune (CMI) response" is one mediated by T-lymphocytes and/or other white blood cells. The CMI immune mechanisms are generally more effective against intracellular infections and disease because the CMI mechanisms prime T cells in a way that, when an antigen appears at a later date, memory T cells are activated to result in a CMI response that destroys target cells that have the corresponding antigen or a portion thereof on their cell surfaces, and thereby the infecting pathogen. The CMI response is focused on the destruction of the source of infection mediated by either effector cells that destroy infected cells of the host by direct cell-to-cell contact and/or by the release of molecules, such as cytokines, that possess anti-viral activity. Thus the CMI response, which is characterised by a specific T lymphocyte cellular response, is crucial to produce resistance to diseases caused by cancer, viruses, pathogenic and other intracellular microorganisms.

T CELLS IMPLICATED IN THE CMI RESPONSE

At least two special types of T cells are required to initiate and/or to enhance CMI and and humoral responses. The antigenic receptors on a particular subset of T cells which express a CD4 co-receptor can be T helper (Th) cells or CD4 T cells (herein after called T helper cells) and they recognise antigenic peptides bound to MHC class II molecules. In contrast, the antigenic receptors on a particular subset of T cells which express a CD8 co-receptor are called Cytotoxic T lymphocytes (CTLs) or CD8+ T cells (hereinafter called CD8+ T cells) and they react with antigens displayed on MHC Class I molecules.

HELPER T CELLS

Helper T cells or CD4+ cells can be further divided into two functionally distinct subsets: Th1 and Th2 which differ in their cytokine and effector function. Th1 and Th2 responses have been shown to be regulated not only in a positive but also in a negative way such that Th1 cellular responses are augmented by Th1 cytokines such as IL-2, IL-12 and IFN-gamma and decreased by Th2 cytokines such as IL-4 and IL-10. In contrast, antibody responses are enhanced by Th2 cytokines such as IL-4 and IL-10 but are downregulated by Th1 cytokines such as IFN-gamma and another cytokine IL-12 that enhances IFN-gamma and is produced by monocytes. Thus, classic Th1 cytokines such as IFN-gamma, IL-2 and IL-12 can be regarded as immune co-factors that induce an effective inflammatory response. In contrast, the classic Th2 cytokines such as IL-4 and IL-10 can be regarded as cytokines that will suppress a severe inflammatory response.

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CD8+ T CELLS

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CD8+ T cells may function in more than one way. The best known function of CD8+ T cells is the killing or lysis of target cells bearing peptide antigen in the context of an MHC class I molecule. Hence the reason why these cells are often termed cytotoxic T lymphocytes (CTL). However, another function, perhaps of greater protective relevance in certain infections is the ability of CD8+ T cells to secrete interferon gamma (IFN-gamma). Thus assays of lytic activity and of IFN-gamma release are both of value in measuring CD8+ T cell immune response (eg in an ELISPOT assay as set forth below). In infectious diseases there is evidence to suggest that CD8+ T cells can protect by killing an infectious agent comprising an infectious antigen at the early stages of a disease before any symptoms of disease are produced.

ENHANCED CMI RESPONSE

The present invention concerns methods, processes and compositions capable of 15 enhancing and/or modulating the CMI response in a host subject against a target antigen. As used herein, the term "enhancing" encompasses improvements in all aspects of the CMI response which include but are not limited to a stimulation and/or augmentation and/or potentiation and/or up-regulation of the magnitude and/or duration, and/or quality of the CMI response to an antigen or a nucleotide sequence 20 encoding an antigen of interest. By way of example, the CMI response may be enhanced by either (i) enhancing the activation and/or production and/or proliferation of CD8+ T cells that recognise a target antigen and/or (ii) shifting the CMI response from a Th2 to a Th1 type response. This enhancement of the Th1 associated responses is of particular value in responding to intracellular infections because, as explained 25 above, the CMI response is enhanced by activated Thl (such as, for example, IFNgamma inducing) cells.

Such an enhanced immune response may be generally characterized by increased titers of interferon-producing CD4⁺ and/or CD8⁺ T lymphocytes, increased antigenspecific CD8+ T cell activity, and a T helper 1-like immune response (Th1) against the antigen of interest (characterized by increased antigen-specific antibody titers of the subclasses typically associated with cellular immunity (such as, for example IgG2a), usually with a concomitant reduction of antibody titers of the subclasses typically associated with humoral immunity (such as, for example IgG1)) instead of a T helper 2-like immune response (Th2).

The enhancement of a CMI response may be determined by a number of well-known assays, such as by lymphoproliferation (lymphocyte activation) assays, CD8+ T cell assays, or by assaying for T-lymphocytes specific for the epitope in a sensitized subject (see, for example, Erickson *et al.* (1993) J. Immunol. 151: 4189-4199; and Doe *et al.* (1994) Eur. J. Immunol. 24: 2369-2376) or CD8+ T cell ELISPOT assays for measuring Interferon gamma production (Miyahara *et al* PNAS(USA) (1998) 95: 3954-3959).

ENHANCED T-CELL RESPONSE

As used herein, the term "enhancing a T -cell response" encompasses improvements in all aspects of the T-cell response which include but are not limited to a stimulation and/or augmentation and/or potentiation and/or up-regulation of the magnitude and/or duration, and/or quality of the T-cell response to an antigen (which may be repeatedly administered) or a nucleotide sequence encoding an antigen. The antigen may be a Chlamydia antigen, preferably a Chlamydia pneumoniae antigen. By way of example, the T-cell response may be enhanced by either enhancing the activation and/or production and/or distribution and/or proliferation of the induced T-cells and/or longevity of the T-cell response to T-cell inducing/modulating antigen or nucleotide sequence encoding an antigen. The enhancement of the T-cell response in a host subject may be associated with the enhancement and/or modulation of the Th1 immune response in the host subject.

The enhancement of the T-cell response may be determined by a number of well-known assays, such as by lymphoproliferation (lymphocyte activation) assays, CD8+T-cell cytotoxic cell assays, or by assaying for T-lymphocytes specific for the epitope in a sensitized subject (see, for example, Erickson et al. (1993) J. Immunol. 151: 4189-4199; and Doe et al. (1994) Eur. J. Immunol. 24: 2369-2376) or CD8+T-cell ELISPOT assays for measuring Interferon gamma production (Miyahara et al PNAS(USA) (1998) 95: 3954-3959).

ANTIGEN

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Each disease causing agent or disease state has associated with it an antigen or immunodominant epitope on the antigen which is crucial in immune recognition and ultimate elimination or control of a disease causing agent or disease state in a host. In order to mount a humoral and/or cellular immune response against a particular disease, the host immune system must come in contact with an antigen or an immunodominant epitope on an antigen associated with that disease state.

As used herein, the term "antigen" refers to any agent, generally a macromolecule, which can elicit an immunological response in an individual. The term "antigen" is used interchangeably with the term "immunogen". The immunological response may be of B- and/or T-lymphocytic cells. The term may be used to refer to an individual macromolecule or to a homogeneous or heterogeneous population of antigenic macromolecules. As used herein, "antigen" is used to refer to a protein molecule or portion thereof which contains one or more antigenic determinants or epitopes. As used herein, the term "antigen" means an immunogenic peptide or protein of interest comprising one or more epitopes capable of inducing a CMI response to an infectious Chlamydia pathogen. The antigen can include but is not limited to an auto-antigen, a self-antigen, a cross-reacting antigen, an alloantigen, a tolerogen, an allergen, a hapten, an immunogen or parts thereof as well as any combinations thereof.

EPITOPE

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As used herein, the term "epitope" generally refers to the site on an antigen which is recognised by a T-cell receptor and/or an antibody. Preferably it is a short peptide derived from or as part of a protein antigen. However the term is also intended to include peptides with glycopeptides and carbohydrate epitopes. Several different epitopes may be carried by a single antigenic molecule. The term "epitope" also includes modified sequences of amino acids or carbohydrates which stimulate

responses which recognise the whole organism. It is advantageous if the selected epitope is an epitope of an infectious agent, such as a *Chlamydia* bacterium, which causes the infectious disease.

SEQ ID Nos 1-76 in the compositions of the invention may be supplemented or substituted with molecules comprising fragments of SEQ ID Nos 1-76. Such fragments may comprise at least n consecutive monomers from the molecules and depending on the particular sequence. n is either (i) 7 or more for protein molecules (eg. 8 18, 20 or more), preferably such that the fragment comprises an epitope from the sequence. or (ii) 10 or more for nucleic acid molecules (eg 15, 18, 20, 25, 30, 35, 40 or more).

SOURCE OF EPITOPES

The epitope can be generated from knowledge the amino acid and corresponding 15 DNA sequences of the peptide or polypeptide, as well as from the nature of particular amino acids (e.g., size, charge, etc.) and the codon dictionary, without undue experimentation. See, e.g., Ivan Roitt, Essential Immunology, 1988; Kendrew, supra; Janis Kuby, Immunology, 1992 e.g., pp. 79-81. Some guidelines in determining whether a protein will stimulate a response, include: Peptide length-preferably the peptide is about 8 or 9 amino acids long to fit into the MHC class I complex and about 20 13-25 amino acids long to fit into a class II MHC complex. This length is a minimum for the peptide to bind to the MHC complex. It is preferred for the peptides to be longer than these lengths because cells may cut peptides. The peptide may contain an appropriate anchor motif which will enable it to bind to the various class I or class II molecules with high enough specificity to generate an immune response (See 25 Bocchia, M. et al, Specific Binding of Leukemia Oncogene Fusion Protein Pentides to HLA Class I Molecules, Blood 85:2680-2684; Englehard, VH, Structure of peptides associated with class I and class II MHC molecules Ann. Rev. Immunol. 12:181 (1994)). This can be done, without undue experimentation, by comparing the sequence of the protein of interest with published structures of peptides associated 30 with the MHC molecules. Thus, the skilled artisan can ascertain an epitope of interest by comparing the protein sequence with sequences listed in the protein data base.

T CELL EPITOPES

Preferably one or more antigens of the present invention contain one or more T cell 35 epitopes. As used herein, the term "T cell epitope" refers generally to those features of a peptide structure which are capable of inducing a T cell response. In this regard, it is accepted in the art that T cell epitopes comprise linear peptide determinants that assume extended conformations within the peptide-binding cleft of MHC molecules (Unanue et al. (1987) Science 236: 551-557). As used herein, a T cell epitope is 40 generally a peptide having at least about 3-5 amino acid residues, and preferably at least 5-10 or more amino acid residues. However, as used herein, the term "T cell epitope" encompasses any MHC Class I-or MHC Class II restricted peptide. The ability of a particular T cell epitope to stimulate/enhance a CMI response may be determined by a number of well-known assays, such as by lymphoproliferation 45 (lymphocyte activation) assays, CD8+ T-cell cytotoxic cell assays, or by assaying for T-lymphocytes specific for the epitope in a sensitized subject. See, e. g., Erickson et al. (1993) J. Immunol. 151: 4189-4199; and Doe et al. (1994) Eur. J. Immunol. 24: 2369-2376 or CD8+ T-cell ELISPOT assays for measuring Interferon gamma production (Miyahara et al PNAS(USA) (1998) 95: 3954-3959). 50

CD8+ T-CELL EPITOPES

Preferably the antigens of the present invention comprisse CD8+ T-cell inducing epitopes. A CD8+ T-cell -inducing epitope is an epitope capable of stimulating the formation, or increasing the activity, of specific CD8+ T-cells following its administration to a host subject. The CD8+ T-cell epitopes may be provided in a variety of different forms such as a recombinant string of one or two or more epitopes. CD8+ T-cell epitopes have been identified and can be found in the literature, for many different diseases. It is possible to design epitope strings to generate CD8+ T-cell response against any chosen antigen that contains such CD8+ T-cell epitopes.

Advantageously, CD8+ T-cell inducing epitopes may be provided in a string of multiple epitopes which are linked together without intervening sequences so that unnecessary nucleic acid material is avoided.

T HELPER EPITOPES

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Preferably the antigens of the present invention comprise helper T lymphocyte epitopes. Various methods are available to identify T helper cell epitopes suitable for use in accordance herewith. For example, the amphipathicity of a peptide sequence is known to effect its ability to function as a T helper cell inducer. A full discussion of T helper cell-inducing epitopes is given in U.S. Patent 5,128,319, incorporated herein by reference.

B CELL EPITOPES

Preferably the antigens of the present invention comprise a mixture of CD8+ T-cell epitopes and B cell epitopes. As used herein, the term "B cell epitope" generally refers to the site on an antigen to which a specific antibody molecule binds. The identification of epitopes which are able to elicit an antibody response is readily accomplished using techniques well known in the art. See, e. g., Geysen et al. (1984) Proc. Natl. Acad. Sci. USA 81: 3998-4002 (general method of rapidly synthesizing peptides to determine the location of immunogenic epitopes in a given antigen); U. S. Patent No. 4,708,871 (procedures for identifying and chemically synthesizing epitopes of antigens); and Geysen et al.(1986) Molecular Immunology 23: 709-715 (technique for identifying peptides with high affinity for a given antibody).

COMBINATION OF EPITOPES

- In a preferred embodiment of the present invention, the antigen or antigen combination comprises a mixture of a CD8+ T-cell -inducing epitopes and a T helper cell-inducing epitopes.
- As is well known in the art, T and B cell inducing epitopes are frequently distinct from each other and can comprise different peptide sequences. Therefore certain regions of a protein's peptide chain can possess either T cell or B cell epitopes. Therefore, in addition to the CD8+ T-cell epitopes, it may be preferable to include one or more epitopes recognised by T helper cells, to augment the immune response generated by the CD8+ T-cell epitopes.

The mechanism of enhancing a CD8+ T-cell induced response in vivo by T helper cell inducing agents is not completely clear. However, without being bound by theory, it is likely that the enhancing agent, by virtue of its ability to induce T helper cells, will result in increased levels of necessary cytokines that assist in the clonal expansion and dissemination of specific CD8+ T-cells. Regardless of the underlying mechanism, it

is envisioned that the use of mixtures of helper T cell and CD8+ T-cell -inducing antigen combinations of the present invention will assist in the enhancement of the CMI response. Particularly suitable T helper cell epitopes are ones which are active in individuals of different HLA types, for example T helper epitopes from tetanus (against which most individuals will already be primed). It may also be useful to include B cell epitopes for stimulating B cell responses and antibody production. Synthetic nucleotide sequences may also be constructed to produce two types of immune responses: T cell only and T cell combined with a B cell response.

10 IMMUNODOMINANT EPITOPE

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When an individual is immunized with an antigen or combination of antigens or nucleotide sequence or combinations of nucleotide sequences encoding multiple epitopes of a target antigen, in many instances the majority of responding T lymphocytes will be specific for one or more linear epitopes from that target antigen and/or a majority of the responding B lymphocytes will be specific for one or more linear or conformational epitopes for the antigen or combination of antigens.. For the purposes of the present invention, then, such epitopes are referred to as "immunodominant epitopes". In an antigen having several immunodominant epitopes, a single epitope may be the most dominant in terms of commanding a specific T or B cell response.

As the Examples show, at least sixteen peptides of the present invention were recognised by IFN-gamma positive CD8+ T cell populations which were actually expanded as a result of bacterial infection.

ADJUVANTS

The compositions of the present invention may be administered in conjunction with other immunoregulatory agents. In particular, the compositions of the present invention may be administered with an adjuvant.

The inclusion of an adjuvant and in particular, a genetic adjuvant may be useful in further enhancing or modulating the CMI response. An adjuvant may enhance the CMI response by enhancing the immunogenicity of a co-administered antigen in an immunized subject, as well inducing a Th1-like immune response against the co-administered antigen which is beneficial in a vaccine product.

An immune response and particularly a CMI response may be refined, by the addition of adjuvants to combinations of antigens or nucleotide sequences encoding combinations of antigens which lead to particularly effective compositions for eliciting a long lived and sustained enhanced CMI response.

As used herein, the term "adjuvant" refers to any material or composition capable of specifically or non-specifically altering, enhancing, directing, redirecting, potentiating or initiating an antigen-specific immune response.

The term "adjuvant" includes but is not limited to a bacterial ADP-ribosylating exotoxin, a biologically active factor, immunomodulatory molecule, biological response modifier or immunostimulatory molecule such as a cytokine, an interleukin, a chemokine or a ligand or an epitope (such as a helper T cell epitope) and optimally combinations thereof which, when administered with an antigen, antigen composition

or nucleotide sequence encoding such antigens enhances or potentiates or modulates the CMI response relative to the CMI response generated upon administration of the antigen or combination of antigens alone. The adjuvant may be any adjuvant known in the art which is appropriate for human or animal use.

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invention.

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Immunomodulatory molecules such as cytokines (TNF-alpha, IL-6, GM-CSF, and IL-2), and co-stimulatory and accessory molecules (B7-1, B7-2) may be used as adjuvants in a variety of combinations. In one embodiment GM-CSF is not administered to subject before, in or after the administration regimen. Simultaneous production of an immunomodulatory molecule and an antigen of interest at the site of 10 expression of the antigen of interest may enhance the generation of specific effectors which may help to enhance the CMI response. The degree of enhancement of the CMI response may be dependent upon the specific immunostimulatory molecules and/or adjuvants used because different immunostimulatory molecules may elicit different mechanisms for enhancing and/or modulating the CMI response. By way of 15 example, the different effector mechanisms/immunomodulatory molecules include but are not limited to augmentation of help signal (IL-2), recruitment of professional APC (GM-CSF), increase in T cell frequency (IL-2), effect on antigen processing pathway and MHC expression (IFN-gamma and TNF-alpha) and diversion of immune response away from the Th1 response and towards a Th2 response (LTB) (see WO 20 97/02045). Unmethylated CpG containing oligonucleotides (see WO96/02555) are also preferential inducers of a Th1 response and are suitable for use in the present

Without being bound by theory, the inclusion of an adjuvant is advantageous because the adjuvant may help to enhance the CMI response to the expressed antigen by diverting the Th2 response to a Th1 response and/or specific effector associated mechanisms to an expressed epitope with the consequent generation and maintenance of an enhanced CMI response (see, for example, the teachings in WO 97/02045).

The inclusion of an adjuvant with an antigen or nucleotide sequence encoding the antigen is also advantageous because it may result in a lower dose or fewer doses of the antigen/antigenic combination being necessary to achieve the desired CMI response in the subject to which the antigen or nucleotide sequence encoding the antigen is administered, or it may result in a qualitatively and/or quantitatively different immune response in the subject. The effectiveness of an adjuvant can be determined by administering the adjuvant with the antigen in parallel with the antigen alone to animals and comparing antibody and/or cellular-mediated immunity in the two groups using standard assays such as radioimmunoassay, ELISAs, CD8+ T-cell assays, and the like, all well known in the art. Typically, the adjuvant is a separate moiety from the antigen, although a single molecule (such for example, CTB) can have both adjuvant and antigen properties.

As used herein, the term "genetic adjuvant" refers to an adjuvant encoded by a nucleotide sequence and which, when administered with the antigen enhances the CMI response relative to the CMI response generated upon administration of the antigen alone.

Bacterial ADP-ribosylating toxins and detoxified derivatives thereof may be used as adjuvants in the invention. Preferably, the protein is derived from E. coli (i.e., E. coli heat labile enterotoxin "LT), cholera ("CT"), or pertussis ("PT"). In one preferred embodiment, the genetic adjuvant is a bacterial ADP-ribosylating

exotoxin.

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ADP-ribosylating bacterial toxins are a family of related bacterial exotoxins and include diphtheria toxin (DT), pertussis toxin (PT), cholera toxin (CT), the E. coli heat-labile toxins (LT1 and LT2), Pseudomonas endotoxin A, Pseudomonas exotoxin 10 S, B. cereus exoenzyme, B. sphaericus toxin, C. botulinum C2 and C3 toxins, C. limosum exoenzyme, as well as toxins from C. perfringens, C. spiriforma and C. difficile, Staphylococcus aureus EDIN, and ADP-ribosylating bacterial toxin mutants such as CRM₁₉₇, a non-toxic diphtheria toxin mutant (see, e.g., Bixler et al. (1989) Adv. Exp. Med. Biol. 251:175; and Constantino et al. (1992) Vaccine). Most ADP-15 ribosylating bacterial toxins are organized as an A:B multimer, wherein the A subunit contains the ADP-ribosyltransferase activity, and the B subunit acts as the binding moiety. Preferred ADP-ribosylating bacterial toxins for use in the compositions of the present invention include cholera toxin and the E. coli heat-labile toxins.

20 Cholera toxin (CT) and the related E. coli heat labile enterotoxins (LT) are secretion products of their respective enterotoxic bacterial strains that are potent immunogens and exhibit strong toxicity when administered systemically, orally, or mucosally. Both CT and LT are known to provide adjuvant effects for antigen when administered via the intramuscular or oral routes. These adjuvant effects have been observed at 25 doses below that required for toxicity. The two toxins are extremely similar molecules, and are at least about 70-80% homologous at the amino acid level.

Preferably the genetic adjuvant is cholera toxin (CT), enterotoxigenic E. Coli heatlabile toxin (LT), or a derivative, subunit, or fragment of CT or LT which retains adjuvanticity. In an even more preferred embodiment, the genetic adjuvant is LT. In 30 another preferred embodiment, the genetic adjuvant may be CTB or LTB.

Preferably the entertoxin is a non-toxic enterotoxin.

The use of detoxified ADP-ribosylating toxins as mucosal adjuvants is described in WO 95/17211 and as parenteral adjuvants in WO 98/42375. The toxin or toxoid is 35 preferably in the form of a holotoxin, comprising both A and B subunits. Preferably, the A subunit contains a detoxifying mutation; preferably the B subunit is not mutated. Preferably, the adjuvant is a detoxified LT mutant such as LT-K63, LT-R72, and LTR192G. The use of ADP-ribosylating toxins and detoxified derivaties thereof, 40 particularly LT-K63 and LT-R72, as adjuvants can be found in the following references each of which is specifically incorporated by reference herein in their entirety (Beignon, et al. Infection and Immunity (2002) 70(6):3012 - 3019; Pizza, et al., Vaccine (2001) 19:2534 - 2541; Pizza, et al., Int. J. Med. Microbiol (2000) 290(4-5):455-461; Scharton-Kersten et al. Infection and Immunity (2000) 68(9):5306 -45 5313; Ryan et al. Infection and Immunity (1999) 67(12):6270 - 6280; Partidos et al. Immunol. Lett. (1999) 67(3):209 - 216; Peppoloni et al. Vaccines (2003) 2(2):285 -293; and Pine et al J. Control Release (2002) 85(1-3):263 - 270). Numerical reference for amino acid substitutions is preferably based on the alignments of the A and B subunits of ADP-ribosylating toxins set forth in Domenighini et al., Mol.

Microbiol (1995) $\underline{15}$ (6):1165 – 1167, specifically incorporated herein by reference in its entirety.

- By way of further example, at least one of the entertoxin subunit coding regions may be genetically modified to detoxify the subunit peptide encoded thereby, for example wherein the truncated A subunit coding region has been genetically modified to disrupt or inactivate ADP-ribosyl transferase activity in the subunit peptide expression product (see, for example, WO 03/004055).
- Thus, these results demonstrate that this genetic adjuvant is particularly desirable where an even more enhanced CMI response is desired. Other desirable genetic adjuvants include but are not limited to nucleotide sequences encoding IL-10, IL-12, IL-13, the interferons (IFNs) (for example, IFN-alpha, IFN-ss, and IFN-gamma), and preferred combinations thereof. Still other such biologically active factors that enhance the CMI response may be readily selected by one of skill in the art, and a suitable plasmid vector containing same constructed by known techniques.

Preferred further adjuvants include, but are not limited to, one or more of the following set forth below:

Mineral Containing Compositions

Mineral containing compositions suitable for use as adjuvants in the invention include mineral salts, such as aluminium salts and calcium salts. The invention includes mineral salts such as hydroxides (e.g. oxyhydroxides), phosphates (e.g. hydroxyphoshpates, orthophosphates), sulphates, etc. {e.g. see chapters 8 & 9 of ref. Bush and Everett (2001) Int J Syst Evol Microbiol 51: 203-220), or mixtures of different mineral compounds, with the compounds taking any suitable form (e.g. gel, crystalline, amorphous, etc.), and with adsorption being preferred. The mineral containing compositions may also be formulated as a particle of metal salt. See WO 00/23105.

Oil-Emulsions

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Oil-emulsion compositions suitable for use as adjuvants in the invention include squalene-water emulsions, such as MF59 (5% Squalene, 0.5% Tween 80, and 0.5% Span 85, formulated into submicron particles using a microfluidizer). See WO 90/14837. Complete Freund's adjuvant (CFA) and incomplete Freund's adjuvant (IFA) may also be used as adjuvants in the invention.

Saponin Formulations

Saponin formulations, may also be used as adjuvants in the invention. Saponins are a heterologous group of sterol glycosides and triterpenoid glycosides that are found in the bark, leaves, stems, roots and even flowers of a wide range of plant species. Saponin from the bark of the *Quillaia saponaria* Molina tree have been widely studied as adjuvants. Saponin can also be commercially obtained from *Smilax ornata* (sarsaprilla), *Gypsophilla paniculata* (brides veil), and *Saponaria officianalis* (soap root). Saponin adjuvant formulations include purified formulations, such as QS21, as well as lipid formulations, such as ISCOMs. Saponin compositions have been purified using High Performance Thin Layer Chromatography (HP-LC) and Reversed Phase High Performance Liquid Chromatography (RP-HPLC). Specific purified fractions using these techniques have been identified, including QS7, QS17, QS18,

QS21, QH-A, QH-B and QH-C. Preferably, the saponin is QS21. A method of production of QS21 is disclosed in U.S. Patent No. 5,057,540. Saponin formulations may also comprise a sterol, such as cholesterol (see WO 96/33739).

Combinations of saponins and cholesterols can be used to form unique particles called Immunostimulating Complexs (ISCOMs). ISCOMs typically also include a phospholipid such as phosphatidylethanolamine or phosphatidyletholine. Any known saponin can be used in ISCOMs. Preferably, the ISCOM includes one or more of Quil A, QHA and QHC. ISCOMs are further described in EP 0 109 942, WO 96/11711 and WO 96/33739. Optionally, the ISCOMS may be devoid of additional detergent. See WO 00/07621.

A review of the development of saponin based adjuvants can be found in Barr et al (1998) Advanced Drug Delivery Reviews 32: 247-271 and Sjolander et al (1998) Advanced Drug Delivery Reviews (1998) 32: 321-338.

Virosomes and Virus Like Particles (VLPs)

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Virosomes and Virus Like Particles (VLPs) can also be used as adjuvants in the invention. These structures generally contain one or more proteins from a virus optionally combined or formulated with a phospholipid. They are generally non-pathogenic, non-replicating and generally do not contain any of the native viral genome. The viral proteins may be recombinantly produced or isolated from whole viruses. These viral proteins suitable for use in virosomes or VLPs include proteins derived from influenza virus (such as HA or NA), Hepatitis B virus (such as core or capsid proteins), Hepatitis E virus, measles virus, Sindbis virus, Rotavirus, Foot-and-Mouth Disease virus, Retrovirus, Norwalk virus, human Papilloma virus, HIV, RNA-

Mouth Disease virus, Retrovirus, Norwalk virus, human Papilloma virus, HIV, RNA-phages, Qß-phage (such as coat proteins), GA-phage, fr-phage, AP205 phage, and Ty (such as retrotransposon Ty protein p1). VLPs are discussed further in WO 03/024480, WO 03/024481; Niikura et al Virology (2002) 293:273 – 280; Lenz et al Journal of Immunology (2001) 5246 – 5355; Pinto, et al Journal of Infectious

Diseases (2003) 188:327 - 338; and Gerber et al Journal of Virology (2001) 75(10):4752 - 4760l; Virosomes are discussed further in, for example, Gluck et al Vaccine (2002) 20:B10-B16.

Bacterial or Microbial Derivatives

Adjuvants suitable for use in the invention include bacterial or microbial derivatives such as:

Non-toxic derivatives of enterobacterial lipopolysaccharide (LPS)

Such derivatives include Monophosphoryl lipid A (MPL) and 3-O-deacylated MPL (3dMPL). 3dMPL is a mixture of 3 De-O-acylated monophosphoryl lipid A with 4, 5 or 6 acylated chains. A preferred "small particle" form of 3 De-O-acylated monophosphoryl lipid A is disclosed in EP 0 689 454. Such "small particles" of 3dMPL are small enough to be sterile filtered through a 0.22 micron membrane (see EP 0 689 454). Other non-toxic LPS derivatives include monophosphoryl lipid A mimics, such as aminoalkyl glucosaminide phosphate derivatives e.g. RC-529. See Johnson et al. (1999) Bioorg Med Chem Lett 9:2273-2278.

Lipid A Derivatives

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Lipid A derivatives include derivatives of lipid A from *Escherichia coli* such as OM-174. OM-174 is described for example in Meraldi *et al.* Vaccine (2003) <u>21</u>:2485 – 2491; Pajak, *et al.* Vaccine (2003) <u>21</u>:836 – 842.

Immunostimulatory oligonucleotides

Immunostimulatory oligonucleotides suitable for use as adjuvants in the invention include nucleotide sequences containing a CpG motif (a sequence containing an unmethylated cytosine followed by guanosine and linked by a phosphate bond). Bacterial double stranded RNA or oligonucleotides containing palindromic or poly(dG) sequences have also been shown to be immunostimulatory.

The CpG's can include nucleotide modifications/analogs such as phosphorothioate modifications and can be double-stranded or single-stranded. Optionally, the guanosine may be replaced with an analog such as 2'-deoxy-7-deazaguanosine. See Kandimalla, et al Nucleic Acids Research (2003) 31(9): 2393 – 2400; WO 02/26757 and WO 99/62923 for examples of possible analog substitutions. The adjuvant effect of CpG oligonucleotides is further discussed in Krieg Nature Medicine (2003) 9(7): 831 – 835; McCluskie, et al FEMS Immunology and Medical Microbiology (2002) 32:179 – 185; WO 98/40100, U.S. Patent No. 6,207,646, U.S. Patent No. 6,239,116, and U.S. Patent No. 6,429,199.

The CpG sequence may be directed to TLR9, such as the motif GTCGTT or TTCGTT. See Kalman et al (1999) (Nature Genetics 21: 385-389). The CpG sequence may be specific for inducing a Th1 immune response, such as a CpG-A ODN, or it may be more specific for inducing a B cell response, such a CpG-B ODN. CpG-A and CpG-B ODNs are discussed in Blackwell, et al J. Immunol. (2003) 170(8):4061 - 4068; Krieg BBRC (2003) 306:948 - 953; and WO 01/95935. Preferably, the CpG is a CpG-A ODN.

Preferably, the CpG oligonucleotide is constructed so that the 5' end is accessible for receptor recognition. Optionally, two CpG oligonucleotide sequences may be attached at their 3' ends to form "immunomers". See, for example, Kandimalla, et al (2003) 31(part 3):664 – 658; Bhagat et al BBRC (2003) 300:853 – 861 and WO 03/035836.

Human Immunomodulators

Human immunomodulators suitable for use as adjuvants in the invention include cytokines, such as interleukins (e.g. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12, etc.), interferons (e.g. interferon-γ), macrophage colony stimulating factor, and tumor necrosis factor.

Bioadhesives and Mucoadhesives

Bioadhesives and mucoadhesives may also be used as adjuvants in the invention. Suitable bioadhesives include esterified hyaluronic acid microspheres (Singh et al. (2001) J. Cont. Rele. 70:267-276) or mucoadhesives such as cross-linked derivatives of poly(acrylic acid), polyvinyl alcohol, polyvinyl pyrollidone, polysaccharides and carboxymethylcellulose. Chitosan and derivatives thereof may also be used as adjuvants in the invention. See for example, WO99/27960.

Microparticles

Microparticles may also be used as adjuvants in the invention. Microparticles (i.e. a particle of ~100nm to ~150 μ m in diameter, more preferably ~200nm to ~30 μ m in diameter, and most preferably ~500nm to ~10 μ m in diameter) formed from materials that are biodegradable and non-toxic (e.g. a poly(o-hydroxy acid), 5 polyhydroxybutyric acid, a polyorthoester, a polyanhydride, a polycaprolactone, etc.), with poly(lactide-co-glycolide) are preferred, optionally treated to have a negativelycharged surface (e.g. with SDS) or a positively-charged surface (e.g. with a cationic detergent, such as CTAB).

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Liposomes

Examples of liposome formulations suitable for use as adjuvants are described in U.S. Patent No. 6,090,406, U.S. Patent No. 5,916,588, and EP 0 626 169.

Polyoxyethylene ether and Polyoxyethylene Ester Formulations 15 Adjuvants suitable for use in the invention include polyoxyethylene ethers and polyoxyethylene esters (WO99/52549). Such formulations further include polyoxyethylene sorbitan ester surfactants in combination with an octoxynol (WO01/21207) as well as polyoxyethylene alkyl ethers or ester surfactants in combination with at least one additional non-ionic surfactant such as an octoxynol 20 (WO01/21152). Preferred polyoxyethylene ethers are selected from the following group: polyoxyethylene-9-lauryl ether (laureth 9), polyoxyethylene-9-steoryl ether, polyoxytheylene-8-steoryl ether, polyoxyethylene-4-lauryl ether, polyoxyethylene-35lauryl ether, and polyoxyethylene-23-lauryl ether.

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Polyphosphazene (PCPP)

PCPP formulations are described, for example, in Andrianov et al Biomaterials (1998) 19(1 - 3):109 - 115; Payne et al Adv. Drug. Delivery Review (1998) 31(3):185-196.

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Muramyl peptides

Examples of muramyl peptides suitable for use as adjuvants in the invention include N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetyl-normuramyl-Lalanyl-D-isoglutamine (nor-MDP), and N-acetylmuramyl-L-alanyl-D-isoglutaminyl-Lalanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphosphoryloxy)-ethylamine MTP-PE).

Imidazoquinolone Compounds

- Examples of imidazoquinolone compounds suitable for use adjuvants in the invention include Imiquamod and its homologues, described further in Stanley, "Imiquimod and 40 the imidazoquinolones: mechanism of action and therapeutic potential" Clin Exp Dermatol (2002) 27(7):571 - 577; and Jones, "Resiquimod 3M", Curr Opin Investig Drugs (2003) 4(2):214 - 218. The invention may also comprise combinations of aspects of one or more of the adjuvants identified above. For example, the following adjuvant compositions may be used in the invention:
- - a saponin and an oil-in-water emulsion (WO99/11241); (1)
 - a saponin (e.g., QS21) + a non-toxic LPS derivative (e.g., 3dMPL) (see WO **(2)** 94/00153);
- a saponin (e.g., QS21) + a non-toxic LPS derivative (e.g., 3dMPL) + a 50 cholesterol;

a saponin (e.g. QS21) + 3dMPL + IL-12 (optionally + a sterol (WO98/57659); combinations of 3dMPL with, for example, QS21 and/or oil-in-water emulsions (European patent applications 0835318, 0735898 and 0761231).

SAF, containing 10% Squalane, 0.4% Tween 80, 5% pluronic-block polymer L121, and thr-MDP, either microfluidized into a submicron emulsion or vortexed to

generate a larger particle size emulsion.

RibiTM adjuvant system (RAS), (Ribi Immunochem) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (Detox™); and

one or more mineral salts (such as an aluminum salt) + a non-toxic derivative of LPS **(7)**

(such as 3dPML).

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Aluminium salts and MF59 are preferred adjuvants for parenteral immunisation. Mutant bacterial toxins are preferred mucosal adjuvants. 15 The composition may include an antibiotic.

Preferably the compositions of the present invention are administered with alum and/or CpG sequences.

Nucleic Acid

The antigens or epitopes of the present invention may be administered as nucleotide sequences encoding the antigens or epitopes. As used herein, the term nucleotide sequence refers to one of more nucleotide sequences which encode one or more epitopes which are used in the compositions or combinations of the present invention. The term "nucleotide sequence (NOI)" is synonymous with the term "polynucleotide" or "nucleic acid". The NOI may be DNA or RNA of genomic or synthetic or of recombinant origin. The NOI may be double-stranded or single-stranded whether representing the sense or antisense strand or combinations thereof. applications, preferably, the NOI is DNA. For some applications, preferably, the NOI is prepared by use of recombinant DNA techniques (e.g. recombinant DNA). For some applications, preferably, the NOI is cDNA. For some applications, preferably, the NOI may be the same as the naturally occurring form.

- The term "nucleic acid" includes DNA and RNA, and also their analogues, such as 35 those containing modified backbones (e.g. phosphorothioates, etc.), and also peptide nucleic acids (PNA), etc. The invention includes nucleic acid comprising sequences complementary to those described above (e.g. for antisense or probing purposes).
- Nucleic acid according to the invention can be prepared in many ways (e.g. by 40 chemical synthesis, from genomic or cDNA libraries, from the organism itself, etc.) and can take various forms (e.g. single stranded, double stranded, vectors, probes, etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other Chlamydial or host cell nucleic acids).

The invention provides a process for producing nucleic acid of the invention, 45 comprising the step of amplifying nucleic acid using a primer-based amplification method (e.g. PCR).

The invention provides a process for producing nucleic acid of the invention, comprising the step of synthesising at least part of the nucleic acid by chemical means.

5 VECTOR

In one embodiment of the present invention, an antigen or antigenic combination or NOI encoding same is administered directly to a host subject. In another embodiment of the present invention, a vector comprising an NOI is administered to a host subject. Preferably the NOI is prepared and/or administered using a genetic vector. As it is well known in the art, a vector is a tool that allows or faciliates the transfer of an 10 entity from one environment to another. In accordance with the present invention, and by way of example, some vectors used in recombinant DNA techniques allow entities, such as a segment of DNA (such as a heterologous DNA segment, such as a heterologous cDNA segment), to be transferred into a host and/or a target cell for the 15 purpose of replicating the vectors comprising the NOI of the present invention and/or expressing the antigens or epitopes of the present invention encoded by the NOI. Examples of vectors used in recombinant DNA techniques include but are not limited to plasmids, chromosomes, artificial chromosomes or viruses. The term "vector" includes expression vectors and/or transformation vectors. The term "expression 20 vector" means a construct capable of in vivo or in vitro/ex vivo expression. The term "transformation vector" means a construct capable of being transferred from one species to another.

NAKED DNA

The vectors comprising the NOI of the present invention may be administered directly as "a naked nucleic acid construct", preferably further comprising flanking sequences homologous to the host cell genome. As used herein, the term "naked DNA" refers to a plasmid comprising the NOI of the present invention together with a short promoter region to control its production. It is called "naked" DNA because the plasmids are not carried in any delivery vehicle. When such a DNA plasmid enters a host cell, such as a eukaryotic cell, the proteins it encodes are transcribed and translated within the cell.

VIRAL VECTORS

Alternatively, the vectors comprising the NOI of the present invention may be introduced into suitable host cells using a variety of viral techniques which are known in the art, such as for example infection with recombinant viral vectors such as retroviruses, herpes simplex viruses and adenoviruses. The vector may be a recombinant viral vectors. Suitable recombinant viral vectors include but are not limited to adenovirus vectors, adeno-associated viral (AAV) vectors, herpes-virus vectors, a retroviral vector, lentiviral vectors, baculoviral vectors, pox viral vectors or parvovirus vectors (see Kestler et al 1999 Human Gene Ther 10(10):1619-32). In the case of viral vectors, administration of the NOI is mediated by viral infection of a target cell.

45 TARGETED VECTOR

The term "targeted vector" refers to a vector whose ability to infect or transfect or transduce a cell or to be expressed in a host and/or target cell is restricted to certain cell types within the host subject, usually cells having a common or similar phenotype.

EXPRESSION VECTOR

Preferably, the NOI of the present invention which is inserted into a vector is operably linked to a control sequence that is capable of providing for the expression of the antigens or epitopes by the host cell, i.e. the vector is an expression vector. The agent produced by a host cell may be secreted or may be contained intracellularly depending on the NOI and/or the vector used. As will be understood by those of skill in the art, expression vectors containing the NOI can be designed with signal sequences which direct secretion of the EOI through a particular prokaryotic or eukaryotic cell membrane.

10 FUSION PROTEINS

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The Chlamydia pneumoniae antigens used in the invention may be present in the composition as individual separate polypeptides, but it is preferred that at least two (i.e. 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20) of the antigens are expressed as a single polypeptide chain (a 'hybrid' polypeptide). Hybrid polypeptides offer two principal advantages: first, a polypeptide that may be unstable or poorly expressed on its own can be assisted by adding a suitable hybrid partner that overcomes the problem; second, commercial manufacture is simplified as only one expression and purification need be employed in order to produce two polypeptides which are both antigenically useful.

The hybrid polypeptide may comprise two or more polypeptide sequences from the first antigen group. Accordingly, the invention includes a composition comprising a first amino acid sequence and a second amino acid sequence, wherein said first and second amino acid sequences are selected from a *Chlamydia* bactgerium, preferably a *Chlamydia pneumoniae* antigen or a fragment thereof of the first antigen group. Preferably, the first and second amino acid sequences in the hybrid polypeptide comprise different epitopes.

The hybrid polypeptide may comprise two or more polypeptide sequences from the second antigen group. Accordingly, the invention includes a composition comprising a first amino acid sequence and a second amino acid sequence, wherein said first and second amino acid sequences are selected from a *Chlamydia pneumoniae* antigen or a fragment thereof of the second antigen group. Preferably, the first and second amino acid sequences in the hybrid polypeptide comprise difference epitopes.

The hybrid polypeptide may comprise one or more polypeptide sequences from the first antigen group and one or more polypeptide sequences from the second antigen group. Accordingly, the invention includes a composition comprising a first amino acid sequence and a second amino acid sequence, said first amino acid sequence selected from a *Chlamydia pneumoniae* antigen or a fragment thereof from the first antigen group and said second amino acid sequence selected from a *Chlamydia* bactgerium, preferably a *Chlamydia pneumoniae* antigen or a fragment thereof from the second antigen group. Preferably, the first and second amino acid sequences in the hybrid polypeptide comprise difference epitopes.

The hybrid polypeptide may comprise one or more polypeptide sequences from the first antigen group and one or more polypeptide sequences from the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group or the ninth antigen group or the tenth antigen group. Accordingly, the invention includes a composition

comprising a first amino acid sequence and a second amino acid sequence, said first amino acid sequence selected from a *Chlamydia pneumoniae* antigen or a fragment thereof from the first antigen group and said second amino acid sequence selected from a *Chlamydia pneumoniae* antigen or a fragment thereof from the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group or the ninth antigen group or the tenth antigen group. Preferably, the first and second amino acid sequences in the hybrid polypeptide comprise difference epitopes.

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- The hybrid polypeptide may comprise one or more polypeptide sequences from the 10 second antigen group and one or more polypeptide sequences from the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group or the ninth antigen group or the tenth antigen group. Accordingly, the invention includes a composition comprising a first amino acid sequence and a second amino acid sequence, said first 15 amino acid sequence selected from a Chlamydia pneumoniae antigen or a fragment thereof from the second antigen group and said second amino acid sequence selected from a Chlamydia pneumoniae antigen or a fragment thereof from the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group or the ninth antigen group or 20 the tenth antigen group. Preferably, the first and second amino acid sequences in the hybrid polypeptide comprise difference epitopes.
- Hybrids consisting of amino acid sequences from two, three, four, five, six, seven, eight, nine, or ten *Chlamydia pneumoniae* antigens are preferred. In particular, hybrids consisting of amino acid sequences from two, three, four, or five *Chlamydia pneumoniae* antigens are preferred. Different hybrid polypeptides may be mixed together in a single formulation. Within such combinations, a *Chlamydia pneumoniae* antigen may be present in more than one hybrid polypeptide and/or as a non-hybrid polypeptide. It is preferred, however, that an antigen is present either as a hybrid or as a non-hybrid, but not as both.

Two-antigen hybrids for use in the invention may comprise any one of the combinations disclosed above.

- Hybrid polypeptides can be represented by the formula NH₂-A-{-X-L-}_n-B-COOH, wherein: X is an amino acid sequence of a *Chlamydia pneumoniae* antigen or a fragment thereof from the first antigen group, the second antigen group or the third antigen group or the fourth antigen group or the fifth antigen group or the sixth antigen group or the seventh antigen group or the eight antigen group or the ninth antigen group or the tenth antigen group.; L is an optional linker amino acid sequence; A is an optional N-terminal amino acid sequence; B is an optional C-terminal amino acid sequence; and n is 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15.
- If a -X- moiety has a leader peptide sequence in its wild-type form, this may be included or omitted in the hybrid protein. In some embodiments, the leader peptides will be deleted except for that of the -X- moiety located at the N-terminus of the hybrid protein *i.e.* the leader peptide of X₁ will be retained, but the leader peptides of X₂ ... X_n will be omitted. This is equivalent to deleting all leader peptides and using the leader peptide of X₁ as moiety -A-.

For each n instances of {-X-L-}, linker amino acid sequence -L- may be present or absent. For instance, when n=2 the hybrid may be NH₂-X₁-L₁-X₂-COOH, NH₂-X₁-X₂-COOH, NH₂-X₁-X₂-COOH, NH₂-X₁-X₂-COOH, etc. Linker amino acid sequence(s) -L- will typically be short (e.g. 20 or fewer amino acids i.e. 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples comprise short peptide sequences which facilitate cloning, poly-glycine linkers (i.e. comprising Gly_n where n = 2, 3, 4, 5, 6, 7, 8, 9, 10 or more), and histidine tags (i.e. His_n where n = 3, 4, 5, 6, 7, 8, 9, 10 or more). Other suitable linker amino acid sequences will be apparent to those skilled in the art. A useful linker is GSGGGG (SEQ ID No 77), with the Gly-Ser dipeptide being formed from a BamHI restriction site, thus aiding cloning and manipulation, and the (Gly)₄ tetrapeptide being a typical poly-glycine linker.

-A- is an optional N-terminal amino acid sequence. This will typically be short (e.g. 40 or fewer amino acids i.e. 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include leader sequences to direct protein trafficking, or short peptide sequences which facilitate cloning or purification (e.g. histidine tags i.e. His, where n = 3, 4, 5, 6, 7, 8, 9, 10 or more). Other suitable N-terminal amino acid sequences will be apparent to those skilled in the art. If X₁ lacks its own N-terminus methionine, -A-is preferably an oligopeptide (e.g. with 1, 2, 3, 4, 5, 6, 7 or 8 amino acids) which provides a N-terminus methionine.

-B- is an optional C-terminal amino acid sequence. This will typically be short (e.g. 40 or fewer amino acids i.e. 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include sequences to direct protein trafficking, short peptide sequences which facilitate cloning or purification (e.g. comprising histidine tags i.e. His, where n = 3, 4, 5, 6, 7, 8, 9, 10 or more), or sequences which enhance protein stability. Other suitable C-terminal amino acid sequences will be apparent to those skilled in the art. Most preferably, n is 2 or 3.

The invention also provides nucleic acid encoding hybrid polypeptides of the invention. Furthermore, the invention provides nucleic acid which can hybridise to this nucleic acid, preferably under "high stringency" conditions (e.g. 65°C in a 0.1xSSC, 0.5% SDS solution).

The NOI of the present invention may be expressed as a fusion protein comprising an adjuvant and/or a biological response modifier and/or immunomodulator fused to the antigens or epitopes of the present invention to further enhance and/or augment the CMI response obtained. The biological response modifier may act as an adjuvant in the sense of providing a generalised stimulation of the CMI response. The antigens or epitopes may be attached to either the amino or carboxy terminus of the biological response modifier.

METHODS OF MAKING

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Polypeptides of the invention can be prepared by various means (e.g. recombinant expression, purification from cell culture, chemical synthesis, etc.) and in various forms (e.g. native, fusions, non-glycosylated, lipidated, etc.). They are preferably prepared in

substantially pure form (i.e. substantially free from other Chlamydial or host cell proteins).

The invention also provides a process for producing a polypeptide of the invention, comprising the step of culturing a host cell transformed with nucleic acid of the invention under conditions which induce polypeptide expression. The invention provides a process for producing a polypeptide of the invention, comprising the step of synthesising at least part of the polypeptide by chemical means. The invention further provides a process for producing a composition according to the invention comprising the step of bringing one or more of SEQ IDs 1-76 into combination with one or more other of SEQ IDs 1-76

Strains

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Preferred polypeptides of the invention comprise an amino acid sequence found in *C.pneumoniae* serovar D, or in one or more of an epidemiologically prevalent serotype. Where hybrid polypeptides are used, the individual antigens within the hybrid (i.e. individual -X- moieties) may be from one or more strains. Where n=2, for instance, X_2 may be from the same strain as X_1 or from a different strain. Where n=3, the strains might be (i) $X_1=X_2=X_3$ (ii) $X_1=X_2\neq X_3$ (iii) $X_1\neq X_2=X_3$ (iv) $X_1\neq X_2\neq X_3$ or (v) $X_1=X_3\neq X_2$, etc.

Heterologous host

Whilst expression of the polypeptides of the invention may take place in *Chlamydia*, the invention preferably utilises a heterologous host. The heterologous host may be prokaryotic (e.g. a bacterium) or eukaryotic. It is preferably *E.coli*, but other suitable hosts include *Bacillus subtilis*, *Vibrio cholerae*, *Salmonella typhi*, *Salmonella typhimurium*, *Neisseria lactamica*, *Neisseria cinerea*, *Mycobacteria* (e.g. *M.tuberculosis*), yeasts, *etc*.

Details as to how the molecules which make up the SEQ IDs 1-76 can be produced and used can be found from the relevant international applications such as WO 00/37494, WO 02/02606 and WO 03/049762 and WO 03/068811 and these details need not be repeated here. Where the composition includes a protein that exists in different nascent and mature forms, the mature form of the protein is preferably used. For example, the mature form of the Chlamydia pneumoniae protein lacking the signal peptide may be used

ADMINISTRATION

Compositions of the invention will generally be administered directly to a patient. Direct delivery may be accomplished by parenteral injection (e.g. subcutaneously, intraperitoneally, intravenously, intramuscularly, or to the interstitial space of a tissue), or by rectal, oral (e.g. tablet, spray), vaginal, topical, transdermal {e.g. see WO99/27961} or transcutaneous {e.g. WO02/074244 and WO02/064162 intranasal {e.g. see WO03/028760} ocular, aural, pulmonary or other mucosal administration. The invention may be used to elicit systemic and/or mucosal immunity.

The compositions of the present invention may be administered, either alone or as part of a composition, via a variety of different routes. Certain routes may be favoured for certain compositions, as resulting in the generation of a more effective immune response, prefereably a CMI response, or as being less likely to induce side effects, or as being easier for administration.

By way of example, the compositions of the present invention may be administered via a systemic route or a mucosal route or a transdermal route or it may be administrered directly into a specific tissue. As used herein, the term "systemic administration" includes but is not limited to any parenteral routes of administration. In particular, parenteral administration includes but is not limited to subcutaneous, intraperitoneal, intravenous, intraarterial, intramuscular, or intrasternal injection, intravenous, intraarterial, or kidney dialytic infusion techniques. Preferably, the systemic, parenteral administration is intramuscular injection.

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In one preferred embodiment of the method, the compositions of the present invention are administered via a transdermal route. While it is believed that any accepted mode and route of immunization can be employed and nevertheless achieve some advantages in accordance herewith, the examples below demonstrate particular advantages with transdermal NOI administration. In this regard, and without being bound by theory, it is believed that transdermal administration of a composition may be preferred because it more efficiently activates the cell mediated immune (CMI) arm of the immune system.

The term "transdermal" delivery intends intradermal (e.g., into the dermis or epidermis), transdermal (e.g., "percutaneous") and transmucosal administration, i.e., delivery by passage of an agent into or through skin or mucosal tissue. See, e.g., Transdermal Drug Delivery: Developmental Issues and Research Initiatives, Hadgraft and Guy (eds.), Marcel Dekker, Inc., (1989); Controlled Drug Delivery: Fundamentals and Applications, Robinson and Lee (eds.), Marcel Dekker Inc., (1987); and Transdermal Delivery of Drugs, Vols. 1-3, Kydonieus and Berner (eds.), CRC Press, (1987). Thus, the term encompasses delivery of an agent using a particle delivery device (e.g., a needleless syringe) such as those described in U.S. Patent No. 5,630,796, as well as delivery using particle-mediated delivery devices such as those described in U.S. Patent No. 5,865,796.

As used herein, the term "mucosal administration" includes but is not limited to oral, intranasal, intravaginal, intrarectal, intratracheal, intestinal and ophthalmic administration.

Mucosal routes, particularly intranasal, intratracheal, and ophthalmic are preferred for protection against natural exposure to environmental pathogens such as RSV, flu virus and cold viruses or to allergens such as grass and ragweed pollens and house dust mites. The enhancement of the immune response, preferably the CMI response will enhance the protective effect against a subsequently encountered target antigen such as an allergen or microbial agent.

In another preferred embodiment of the present invention, the compositions of the present invention may be administered to cells which have been isolated from the host subject. In this preferred embodiment, preferably the composition is administered to professional antigen presenting cells (APCs), such as dendritic cells. APCs may be derived from a host subject and modified *ex vivo* to express an antigen of interest and then transferred back into the host subject to induce an enhanced CMI response. Dendritic cells are believed to be the most potent APCs for stimulating enhanced CMI responses because the expressed epitopes of the antigen of interest must be acquired, processed and presented by professional APCs to T cells (both Th1 and Th2 helper cells as well as CD8+ T-cells) in order to induce an enhanced CMI response.

PARTICLE ADMINISTRATION

Particle-mediated methods for delivering the compositions of the present invention are known in the art. Thus, once prepared and suitably purified, the above-described antigens or NOI encoding same can be coated onto core carrier particles using a variety of techniques known in the art. Carrier particles are selected from materials which have a suitable density in the range of particle sizes typically used for intracellular delivery from a gene gun device. The optimum carrier particle size will, of course, depend on the diameter of the target cells.

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By "core carrier" is meant a carrier on which a guest antigen or guest nucleic acid (e.g., DNA, RNA) is coated in order to impart a defined particle size as well as a sufficiently high density to achieve the momentum required for cell membrane penetration, such that the guest molecule can be delivered using particle-mediated techniques (see, e.g., U.S. Patent No. 5,100,792). Core carriers typically include materials such as tungsten, gold, platinum, ferrite, polystyrene and latex. See e.g., Particle Bombardment Technology for Gene Transfer, (1994) Yang, N. ed., Oxford University Press, New York, NY pages 10-11. Tungsten and gold particles are preferred. Tungsten particles are readily available in average sizes of 0.5 to 2.0 microns in diameter. Gold particles or microcrystalline gold (e. g., gold powder A1570, available from Engelhard Corp., East Newark, NJ) will also find use with the present invention. Gold particles provide uniformity in size (available from Alpha Chemicals in particle sizes of 1-3 microns, or available from Degussa, South Plainfield, NJ in a range of particle sizes including 0.95 microns). Microcrystalline gold provides a diverse particle size distribution, typically in the range of 0.5-5 microns. However, the irregular surface area of microcrystalline gold provides for highly efficient coating with nucleic acids. A number of methods are known and have been described for coating or precipitating NOIs onto gold or tungsten particles. Most such methods generally combine a predetermined amount of gold or tungsten with plasmid DNA, CaCl2 and spermidine. The resulting solution is vortexed continually during the coating procedure to ensure uniformity of the reaction mixture. After precipitation of the NOI, the coated particles can be transferred to suitable membranes and allowed to dry prior to use, coated onto surfaces of a sample module or cassette, or loaded into a delivery cassette for use in particular gene gun instruments.

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The particle compositions or coated particles are administered to the individual in a manner compatible with the dosage formulation, and in an amount that will be effective for the purposes of the invention. The amount of the composition to be delivered (e. g., about 0.1 mg to 1 mg, more preferably 1 to 50 ug of the antigen or allergen, depends on the individual to be tested. The exact amount necessary will vary depending on the age and general condition of the individual to be treated, and an appropriate effective amount can be readily determined by one of skill in the art upon reading the instant specification.

45 HOST MAMMALIAN SUBJECT

As used herein, the term "host mammalian subject" means any member of the subphylum cordata, including, without limitation, humans and other primates, including non-human primates such as chimpanzees and other apes and monkey species; farm animals such as cattle, sheep, pigs, goats and horses; domestic mammals such as dogs and cats; laboratory animals including rodents such as mice, rats and

guinea pigs; birds, including domestic, wild and game birds such as chickens, turkeys and other gallinaceous birds, ducks, geese, and the like. The terms do not denote a particular age. Thus, both adult and newborn individuals are intended to be covered. The methods described herein are intended for use in any of the above vertebrate species, since the immune systems of all of these vertebrates operate similarly. If a mammal, the subject will preferably be a human, but may also be a domestic livestock, laboratory subject or pet animal.

The mammal is preferably a human. Where the vaccine is for prophylactic use, the human is preferably a child (e.g. a toddler or infant) or a teenager; where the vaccine is for therapeutic use, the human is preferably a teenager or an adult. A vaccine intended for children may also be administered to adults e.g. to assess safety, dosage, immunogenicity, etc.

PREVENT AND/OR TREAT

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The invention also provides the use of the compositions of the invention in the manufacture of a medicament for raising an immune response in a mammal. The medicament is preferably a vaccine and to the preparation of a vaccine to prevent and/or treat an disorder associated with a *Chlamydia* bacterium. It is to be appreciated that all references herein to treatment include curative, palliative and prophylactic treatment.

The administration of antigenic combinations of the present invention or a composition comprising the NOI encoding the antigenic combinations may be for either "prophylactic" or "therapeutic" purpose. As used herein, the term "therapeutic" or "treatment" includes any of following: the prevention of infection or reinfection; the reduction or elimination of symptoms; and the reduction or complete elimination of a pathogen. Treatment may be effected prophylactically (prior to infection) or therapeutically (following infection).

Prophylaxis or therapy includes but is not limited to eliciting an effective immune response, preferably a CMI immune response and/or alleviating, reducing, curing or at least partially arresting symptoms and/or complications resulting from a T cell mediated immune disorder. When provided prophylactically, the composition of the present invention is typically provided in advance of any symptom. The prophylactic administration of the composition of the present invention is to prevent or ameliorate any subsequent infection or disease. When provided therapeutically, the composition of the present invention is typically provided at (or shortly after) the onset of a symptom of infection or disease. Thus the composition of the present invention may be provided either prior to the anticipated exposure to a disease causing agent or disease state or after the initiation of an infection or disease.

Whether prophylactic or therapeutic administration (either alone or as part of a composition) is the more appropriate will usually depend upon the nature of the disease. By way of example, immunotherapeutic composition of the present invention could be used in immunotherapy protocols to actively inducing immunity by vaccination. This latter form of treatment is advantageous because the immunity is prolonged. On the other hand a vaccine composition will preferably, though not necessarily be used prophylactically to induce an effective CMI response against subsequently encountered antigens or portions thereof (such as epitopes) related to the target antigen.

These uses and methods are preferably for the prevention and/or treatment of a disease caused by a *Chlamydia* (e.g. trachoma, pelvic inflammatory disease, epididymitis, infant pneumonia, artherosclerosis, cardiovascular disease etc.). The compositions may also be effective against *C.pneumoniae*.

PROPHYLACTICALLY OR THERAPEUTICALLY OR IMMUNOLOGICALLY EFFECTIVE AMOUNT

The composition dose administrated to a host subject, in the context of the present invention, should be sufficient to effect a beneficial prophylactic or therapeutic immune response, preferably a CMI response in the subject over time.

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The invention also provides a method for raising an immune response in a mammal comprising the step of administering an effective amount of a composition of the invention. The immune response is preferably protective and preferably involves antibodies and/or cell-mediated immunity. The method may raise a booster response.

As used herein, the term ""prophylactically or therapeutically effective dose" means a dose in an amount sufficient to elicit an enhanced immune response, preferably a CMI response to one or more antigens or epitopes and/or to alleviate, reduce, cure or at least partially arrest symptoms and/or complications from a T cell mediated immune disorder.

Immunogenic compositions used as vaccines comprise an immunologically effective amount of antigen(s), as well as any other components, as needed. By 'immunologically effective amount', it is meant that the administration of that amount to an individual, either in a single dose or as part of a series, is effective for treatment or prevention. This amount varies depending upon the health and physical condition of the individual to be treated, age, the taxonomic group of individual to be treated (e.g. non-human primate, primate, etc.), the capacity of the individual's immune system to synthesise antibodies, the degree of protection desired, the formulation of the vaccine, the treating doctor's assessment of the medical situation, and other relevant factors. It is expected that the amount will fall in a relatively broad range that can be determined through routine trials.

One way of checking efficacy of therapeutic treatment involves monitoring *Chlamydia* infection after administration of the composition of the invention. One way of checking efficacy of prophylactic treatment involves monitoring immune responses against the *Chlamydia antigen*, such as the *Chlamydia pneumoniae* antigen in the compositions of the invention after administration of the composition.

The vaccine compositions of the present invention can be evaluated in *in vitro* and *in vivo* animal models prior to host, e.g., human, administration. For example, *in vitro* neutralization by Peterson *et al* (1988) is suitable for testing vaccine compositions directed toward *Chlamydia*, preferably *Chlamydia pneumoniae*.

One example of such an *in vitro* test is described as follows. Hyper-immune antisera is diluted in PBS containing 5% guinea pig serum, as a complement source. *Chlamydia pneumoniae* (10⁴ IFU; inclusion forming units) are added to the antisera dilutions. The antigen-antibody mixtures are incubated at 37°C for 45 minutes and

inoculated into duplicate confluent Hep-2 or HeLa cell monolayers contained in glass vials (e.g., 15 by 45 mm), which have been washed twice with PBS prior to inoculation. The monolayer cells are infected by centrifugation at 1000X g for 1 hour followed by stationary incubation at 37°C for 1 hour. Infected monolayers are incubated for 48 or 72 hours, fixed and stained with *Chlamydia* specific antibody, such as anti-MOMP. Inclusion-bearing cells are counted in ten fields at a magnification of 200X. Neutralization titer is assigned on the dilution that gives 50% inhibition as compared to control monolayers/IFU.

10 The efficacy of vaccine compositions can also be determined in vivo by challenging animal models of Chlamydia pneumoniae infection, e.g., guinea pigs or mice, with the vaccine compositions. For example, in vivo vaccine composition challenge studies in the guinea pig model of Chlamydia pneumoniae infection can be performed. description of one example of this type of approach follows. Female guinea pigs weighing 450 - 500 g are housed in an environmentally controlled room with a 12 15 hour light-dark cycle and immunized with vaccine compositions via a variety of immunization routes. Post-vaccination, guinea pigs are infected in the genital tract with the agent of guinea pig inclusion conjunctivitis (GPIC), which has been grown in HeLa or McCoy cells (Rank et al. (1988)). Each animal receives approximately 1.4x10⁷ inclusion forming units (IFU) contained in 0.05 ml of sucrose-phosphate-20 glutamate buffer, pH 7.4 (Schacter, 1980). The course of infection monitored by determining the percentage of inclusion-bearing cells by indirect immunofluorescence with GPIC specific antisera, or by Giemsa-stained smear from a scraping from the genital tract (Rank et al 1988). Antibody titers in the serum is determined by an 25 enzyme-linked immunosorbent assay.

Alternatively, in vivo vaccine compositions challenge studies can be performed in the murine model of Chlamydia pneumoniae (Morrison et al 1995). A description of one example of this type of approach is as follows. Female mice 7 to 12 weeks of age receive 2.5 mg of depoprovera subcutaneously at 10 and 3 days before vaginal infection. Post-vaccination, mice are infected in the genital tract with 1,500 inclusion-forming units of Chlamydia pneumoniae contained in 5ml of sucrose-phosphate-glutamate buffer, pH 7.4. The course of infection is monitored by determining the percentage of inclusion-bearing cells by indirect immunofluorescence with Chlamydia pneumoniae specific antisera, or by a Giemsa-stained smear from a scraping from the genital tract of an infected mouse. The presence of antibody titers in the serum of a mouse is determined by an enzyme-linked immunosorbent assay

DOSAGE

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Prophylaxis or therapy can be accomplished by a single direct administration at a single time point or multiple time points. Administration can also be delivered to a single or to multiple sites. Some routes of administration, such as mucosal administration via ophthalmic drops may require a higher dose. Those skilled in the art can adjust the dosage and concentration to suit the particular route of delivery.

Dosage treatment can be a single dose schedule or a multiple dose schedule. multiple doses may be used in a primary immunisation schedule and/or in a booster immunisation schedule. in a multiple dose schedule the various doses may be given by the same or different routes e.g. a parenteral prime and mucosal boost, a mucosal prime and parenteral boost, etc.

HOMOLOGUES

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SEQ IDs 1-76 in the compositions of the invention may be supplemented or substituted with molecules comprising sequences homologous (ie. sharing sequence identity) to SEQ ID Nos 1-76.

Proteins (including protein antigens) as used in the invention (as encoded by the NOI) may have homology and/or sequence identity with naturally occurring forms. Similarly coding sequences capable of expressing such proteins will generally have homology and/or sequence identity with naturally occurring sequences. Techniques for determining nucleic acid and amino acid "sequence identity" also are known in the art. Typically, such techniques include determining the nucleotide sequence of the mRNA for a gene and/or determining the amino acid sequence encoded thereby, and comparing these sequences to a second nucleotide or amino acid sequence.

In general, "identity" refers to an exact nucleotide-to-nucleotide or amino acid-to-amino acid correspondence of two polynucleotides or polypeptide sequences, respectively. Two or more sequences (polynucleotide or amino acid) can be compared by determining their "percent identity." The percent identity of two sequences, whether nucleic acid or amino acid sequences, is the number of exact matches between two aligned sequences divided by the length of the shorter sequences and multiplied by 100.

An approximate alignment for nucleic acid sequences is provided by the local 30 homology algorithm of Smith and Waterman, Advances in Applied Mathematics 2: 482-489 (1981). This algorithm can be applied to amino acid sequences by using the scoring matrix developed by Dayhoff, Atlas of Protein Sequences and Structure, M. O. Dayhoff ed., 5 suppl. 3: 353-358, National Biomedical ResearchFoundation, Washington, D. C., USA, and normalized by Gribskov, Nucl. AcidsRes. 14 (6): 6745-6763 (1986). An exemplary implementation of this algorithm to determine percent 35 identity of a sequence is provided by the Genetics Computer Group (Madison, WI) in the "BestFit" utility application. The default parameters for this method are described in the Wisconsin Sequence Analysis Package Program Manual, Version 8 (1995) (available from Genetics Computer 40 Madison, WI). A preferred method of establishing percent identity in the context of the present invention is to use the MPSRCH package of programs copyrighted by the University of Edinburgh, developed by John F. Collins and Shane S.

Sturrok, and distributed by IntelliGenetics, Inc. (Mountain View, CA). From this suite of packages the Smith-Waterman algorithm can be employed where default parameters are used for the scoring table (for example, gap open penalty of 12, gap extension penalty of one, and a gap of six). From the data generated the "Match" value reflects sequence identity." Other suitable programs for calculating the percent identity or similarity between sequences are generally known in the art, for example, another alignment program is BLAST, used with default parameters. For example,

BLASTN and BLASTP can be used using the following default parameters: genetic code = standard; filter = none; strand = both; cutoff= 60; expect = 10; Matrix = BLOSUM62; Descriptions = 50 sequences; sort by = HIGH SCORE; Databases = non-redundant, GenBank +EMBL + DDBJ + PDB + GenBank CDS translations + Swiss protein + Spupdate + PIR. Details of these programs can be found at the following internet address: http://www.ncbi.nlm.gov/cgi-bin/BLAST.

Alternatively, homology can be determined by hybridization of polynucleotides under conditions which form stable duplexes between homologous regions, followed by digestion with single-stranded-specific nuclease (s), and size determination of the digested fragments. Two DNA, or two polypeptide sequences are "substantially homologous" to each other when the sequences exhibit at least about 80%-85%, preferably at least about 90%, and most preferably at least about 95%-98% sequence identity over a defined length of the molecules, as determined using the methods above.

As used herein, substantially homologous or homologous also refers to sequences showing complete identity to the specified DNA or polypeptide sequence. DNA sequences that are substantially homologous or homologous can be identified in a Southern hybridization experiment under, for example, stringent conditions, as defined for that particular system. For example, stringent hybridization conditions can include 50% formamide, 5x Denhardt's Solution, 5x SSC, 0.1% SDS and 100 pg/ml denatured salmon sperm DNA and the washing conditions can include 2x SSC, 0.1% SDS at 37 C followed by lx SSC, 0.1% SDS at 68 C. Defining appropriate hybridization conditions is within the skill of the art.

Preferably the degree of identity is preferably greater than 50% (eg. 65%. 80%. 90%. or more) and include mutants and allelic variants. Sequence identity between the proteins is preferably determined by the Smith-Waterman homology search algorithm as implemented in the MPSRCH program (Oxford. Molecular). using an affine gap search with parameters gap open penalty=12 and gap extension penalty=1.

SEQ IDs 1-76 in the compositions of the invention may be supplemented or substituted with nucleic acid which can hybridise to the *Chlamydia* nucleic acid. preferably underv"high stringency"conditionsv(c. 65 C in an 0.1 x SSC, 0.5% SDS solution).

Hypothetical Protein

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As used herein, the term "hypothetical protein" refers to a protein which lacks a known cellular location or a known cellular function. Typically, a hypothetical protein lacks significant homologies with known well characterised proteins.

COMPOSITIONS

The invention also provides the compositions of the invention for use as medicaments

(eg. as immunogenic compositions or vaccines) or as diagnostic reagents for detecting a Chylamydia infectioin in a host subject. It also provides the use of the compositions in the manufacture of: (i) a medicament for treating or preventing infection due to Chlamydia pneumoniae bacteria: (ii) a diagnostic reagent for detecting the presence of Chlamydia Pneumonaie bacteria or of antibodies raised against Chlamydia

Pneumonaie bacteria; and/or (iii) a reagent which can raise antibodies against

Chlamydia

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bacteria.

The invention also provides a method of treating a patient, comprising administering to the patient a therapeutically effective amount of a composition according to the invention.

The present invention provides compositions that are useful for preventing and/or treating T cell mediated immune disorders. In one embodiment, the composition is a pharmaceutical composition. In another preferred embodiment, the composition is an immunotherapeutic composition. In an even more preferred embodiment, the composition is a vaccine composition. The composition may also comprise a carrier such as a pharmaceutically or immunologically acceptable carrier. Pharmaceutically acceptable carriers or immunologically acceptable carriers are determined in part by the particular composition being administered as well as by the particular method used to administer the composition. Accordingly, there is a wide variety of suitable formulations of pharmaceutical compositions or vaccine compositions or immunotherapeutic compositions of the present invention.

Immunogenic compositions and medicaments

Compositions of the invention are preferably immunogenic compositions, and are more preferably vaccine compositions. The pH of the composition is preferably between 6 and 8, preferably about 7. The pH may be maintained by the use of a buffer. The composition may be sterile and/or pyrogen-free. The composition may be isotonic with respect to humans.

Vaccines according to the invention may either be prophylactic (i.e. to prevent infection) or therapeutic (i.e. to treat infection), but will typically be prophylactic. Accordingly, the invention includes a method for the therapeutic or prophylactic treatment of Chlamydia pneumoniae infection in an animal susceptible to Chlamydial infection comprising administering to said animal a therapeutic or prophylactic amount of the immunogenic compositions of the invention. Preferably, the immunogenic composition comprises a combination of Chlamydia pneumoniae antigens, said combination selected from the group consisting of two, three, four, five or all six Chlamydia pneumoniae antigens of the first antigen group. Still more preferably, the combination consists of all six Chlamydia pneumoniae antigens of the first antigen group.

Alternatively, the immunogenic composition comprises a combination of *Chlamydia pneumoniae* antigens, said combination selected from the group consisting of two, three, four, five, six, seven, eight, nine, ten, eleven, or twelve *Chlamydia pneumoniae* antigens selected from the first antigen group and the second antigen group. Preferably, the combination is selected from the group consisting of three, four, or five *Chlamydia pneumoniae* antigens selected from the second antigen group. Still more preferably, the combination consists of five *Chlamydia pneumoniae* antigens selected from the second antigen group.

Alternatively, the immunogenic composition comprises a combination of *Chlamydia pneumoniae* antigens, said combination consisting of two, three, four, or five *Chlamydia pneumoniae* antigens of the first antigen group and one, two, three, four, five or six *Chlamydia pneumoniae* antigens of the third antigen group. Preferably, the

combination consists of three, four or five *Chlamydia pneumoniae* antigens of the first antigen group and one, two, three, four, five or six *Chlamydia pneumoniae* antigens of the third antigen group.

Alternatively, the immunigenic composition comprises a combination of Chlamydia pneumoniae antigens, said combination consisting of two, three, four, five, six, seven, eight, nine, ten, eleven or twelve Chlamydia pneumoniae antigens of the first antigen group and the second antigen group and one, two, three, four, five or six Chlamydia pneumoniae antigens of the third antigen group. Preferably, the combination is selected from the group consisting of three, four, or five Chlamydia pneumoniae antigens from the second antigen group and three, four or five Chlamydia pneumoniae from the third antigen group: Still more preferably, the combination consists of five Chlamydia pneumoniae antigens from the second antigen group and three, four or five Chlamydia pneumoniae antigens of the third antigen group and three, four or five Chlamydia pneumoniae antigens of the third antigen group.

In certain embodiments, the composition comprises molecules from different *Chlamydia* species. In some embodiments, the composition may comprise molecules from different serogroups and/or strains of the same *Chlamydia* species. Further embodiments comprise mixtures of one or more *Chlamydia* molecules from different strains.

Many proteins are relatively conserved between different species serogroups and strains of Chlamydia trachomatis and Chlamydia pneumoniae. To ensure maximum cross-strain recognition and reactivity, regions of proteins that are conserved between different Chlamydia species, serogroups and strains can be used in the compositions of the present invention. The invention therefore provides proteins which comprise stretches of amino acid sequence that are shared across the majority of Chlamydia strains. Preferably, therefore, the composition comprises a protein comprising a fragment of a Chlamydia pneumoniae protein (preferably a protein from SEQ ID Nos 1-76 or more preferably SEQ ID Nos 1-41 wherein said fragment consists of n consecutive conserved amino acids.

Further antigens

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The compositions of the invention may further comprise antigen derived from one or more sexually transmitted diseases in addition to *Chlamydia trachomatis*. Preferably the antigen is derived from one or more of the following sexually transmitted diseases: *N.gonorrhoeae* {e.g. i, ii, iii, iv}; human papiloma virus; *Treponema pallidum*; herpes simplex virus (HSV-1 or HSV-2); HIV (HIV-1 or HIV-2); and *Haemophilus ducreyi*.

A preferred composition comprises: (1) at least t of the Chlamydia pneumoniae antigens from either the first antigen group or the second antigen group, where t is 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12 or 13, preferably t is five; (2) one or more antigens from another sexually transmitted disease. Preferably, the sexually transmitted disease is selected from the group consisting of herpes simplex virus, preferably HSV-1 and/or HSV-2; human papillomavirus; N.gonorrhoeae; Treponema pallidum; and Haemophilus ducreyi. These compositions can thus provide protection against the following sexually-transmitted diseases: Chlamydia, genital herpes, genital warts, gonorrhoea, syphilis and chancroid (see Stephens et al (1998) Science 282: 754-759).

Where a saccharide or carbohydrate antigen is used, it is preferably conjugated to a carrier protein in order to enhance immunogenicity (For example, Ramsay et al. (2001) Lancet 357(9251):195-196; Lindberg (1999) Vaccine 17 Suppl 2:S28-36; Buttery & Moxon (2000) JR Coll Physicians Lond 34:163-168; Ahmad & Chapnick (1999) Infect Dis Clin North Am 13:113-133; Goldblatt (1998) J. Med. Microbiol. 47:563-567; European patent 0 477 508; US Patent No. 5,306,492; International patent application WO98/42721; Conjugate Vaccines (eds. Cruse et al.) ISBN 3805549326, particularly vol. 10:48-114; and Hermanson (1996) Bioconjugate Techniques ISBN: 0123423368 or 012342335).

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Preferred carrier proteins are bacterial toxins or toxoids, such as diphtheria or tetanus toxoids. The CRM₁₉₇ diphtheria toxoid is particularly preferred (*Research Disclosure*, 453077 (Jan 2002). Other carrier polypeptides include the *N.meningitidis* outer membrane protein EP-A-0372501), synthetic peptides (EP-A-0378881, EP-A-0427347), heat shock proteins (WO93/17712, WO94/03208) pertussis proteins (WO98/58668, EP-A-0471177) protein D from *H.influenzae* (WO00/56360) cytokines (WO91/01146), lymphokines, hormones, growth factors, toxin A or B from *C.difficile* (WO00/61761) iron-uptake proteins WO01/72337) etc. Where a mixture comprises capsular saccharides from both serogroups A and C, it may be preferred that the ratio (w/w) of MenA saccharide:MenC saccharide is greater than 1 (e.g. 2:1, 3:1, 4:1, 5:1, 10:1 or higher). Different saccharides can be conjugated to the same or different type of carrier protein. Any suitable conjugation reaction can be used, with any suitable linker where necessary.

Toxic protein antigens may be detoxified where necessary e.g. detoxification of pertussis toxin by chemical and/or genetic means. Where a diphtheria antigen is included in the composition it is preferred also to include tetanus antigen and pertussis antigens. Similarly, where a tetanus antigen is included it is preferred also to include diphtheria and pertussis antigens. Similarly, where a pertussis antigen is included it is preferred also to include diphtheria and tetanus antigens.

Antigens in the composition will typically be present at a concentration of at least 1µg/ml each. In general, the concentration of any given antigen will be sufficient to elicit an immune response against that antigen. As an alternative to using protein antigens in the composition of the invention, nucleic acid encoding the antigen may be used Robinson & Torres (1997) Seminars in Immunology 9:271-283; Donnelly et al. (1997) Annu Rev Immunol 15:617-648; Scott-Taylor & Dalgleish (2000) Expert Opin Investig Drugs 9:471-480; Apostolopoulos & Plebanski (2000) Curr Opin Mol Ther 2:441-447; Ilan (1999) Curr Opin Mol Ther 1:116-120; Dubensky et al. (2000) Mol Med 6:723-732; Robinson & Pertmer (2000) Adv Virus Res 55:1-74; Donnelly et al. (2000) Am J Respir Crit Care Med 162(4 Pt 2):S190-193 and Davis (1999) Mt. Sinai J. Med. 66:84-90. Protein components of the compositions of the invention may thus be replaced by nucleic acid (preferably DNA e.g. in the form of a plasmid) that encodes the protein.

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DISEASE STATES

The compositions of the present invention may be used to prevent and/or treat disorders such as but not limited to: pneumonia, cardiovascular diseases, atherosclerosis, bronchitis, pharyngitis, laryngitis, sinusitis, obstructive lung diseases, asthma, chronic obstructive pulmonary disease, reactive arthritis, otitis media,

abdominal aortic aneurysm, erythema nodosum, Reiter syndrome, sarcoidosis, Alzheimer's disease, multiple sclerosis, lymphogranuloma venereum, ocular trachoma, pelvic inflammatory disease, inclusion conjunctivitis, genital trachoma, infant pneumonitis, incipient trachoma, keratitis, papillary hypertrophy, corneal infiltration, vulvovaginitis, mucopurulent rhinitis, salpingitis, cervicitis, cervical follicles, prostatitis, proctitis, urethritis, lymphogranule inguinale, climatic bubo, tropical bubo, and/oresthiomene.

FORMULATIONS

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10 Chlamydial infections affect various areas of the body and so the compositions of the invention may be prepared in various forms. For example, the compositions may be prepared as injectables, either as liquid solutions or suspensions. Solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared (e.g. a lyophilised composition). The composition may be prepared for topical administration e.g. as an ointment, cream or powder. The composition may be 15 prepared for oral administration e.g. as a tablet or capsule, as a spray, or as a syrup (optionally flavoured). The composition may be prepared for pulmonary administration e.g. as an inhaler, using a fine powder or a spray. The composition may be prepared as a suppository or pessary. The composition may be prepared for nasal, aural or ocular administration e.g. as drops. The composition may be in kit form, 20 designed such that a combined composition is reconstituted just prior to administration to a patient. Such kits may comprise one or more antigens in liquid form and one or more lyophilised antigens.

25 Further components of the composition -

The composition of the invention will typically, in addition to the components mentioned above, comprise one or more 'pharmaceutically acceptable carriers', which include any carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition. Suitable carriers are typically large, slowly metabolised macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and lipid aggregates (such as oil droplets or liposomes). Such carriers are well known to those of ordinary skill in the art. The vaccines may also contain diluents, such as water, saline, glycerol, etc. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present. A thorough discussion of pharmaceutically acceptable excipients is available in Gennaro (2000) Remington: The Science and Practice of Pharmacy. 20th ed., ISBN: 0683306472.

The biological molecules of the present invention be formulated into a pharmaceutical composition or an immunotherapeutic composition or a vaccine composition. Such formulations comprise biological molecules combined with a pharmaceutically acceptable carrier, such as sterile water or sterile isotonic saline. Such formulations may be prepared, packaged, or sold in a form suitable for bolus administration or for continuous administration. Injectable formulations may be prepared, packaged, or sold in unit dosage form, such as in ampoules or in multi-dose containers containing a preservative. Formulations include, but are not limited to, suspensions, solutions, emulsions in oily or aqueous vehicles, pastes, and implantable sustained-release or biodegradable formulations. Such formulations may further comprise one or more additional ingredients including, but not limited to, suspending, stabilizing, or dispersing agents. In one embodiment of a formulation for parenteral administration,

the active ingredient is provided in dry (for eg, a powder or granules) form for reconstitution with a suitable vehicle (e. g., sterile pyrogen-free water) prior to parenteral administration of the reconstituted composition. The pharmaceutical compositions may be prepared, packaged, or sold in the form of a sterile injectable aqueous or oily suspension or solution. This suspension or solution may be formulated according to the known art, and may comprise, in addition to the active ingredient, additional ingredients such as the dispersing agents, wetting agents, or suspending agents described herein. Such sterile injectable formulations may be prepared using a non-toxic parenterally-acceptable diluent or solvent, such as water or 1,3-butane diol, for example. Other acceptable diluents and solvents include, but are not limited to, Ringer's solution, isotonic sodium chloride solution, and fixed oils such as synthetic mono-or di-glycerides. Other parentally-administrable formulations which are useful include those which comprise the active ingredient in microcrystalline form, in a liposomal preparation, or as a component of a biodegradable polymer systems. Compositions for sustained release or implantation may comprise pharmaceutically acceptable polymeric or hydrophobic materials such as an emulsion, an ion exchange resin, a sparingly soluble polymer, or a sparingly soluble salt.

KITS

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Also included in the invention is a kit for enhancing a CMI response to the biological 20 molecules of the present invention. Such a kit may comprise an antigenic composition or nucleotide sequence encoding same. The kit may also include an adjuvant, preferably a genetic adjuvant is administered with or as part of the biological molecule and instructions for administering the biological molecule. Other preferred components of the kit include an applicator for administering the biological molecule. 25 As used herein, the term "applicator" refers to any device including but not limited to a hypodermic syringe, gene gun, particle acceleration device, nebulizer, dropper, bronchoscope, suppository, impregnated or coated vaginally-insertable material such as a tampon, douche preparation, solution for vaginal irrigation, retention enema preparation, suppository, or solution for rectal or colonic irrigation for applying the 30 NOI either systemically or mucosally or transdermally to the host subject.

The invention also provides for a kit comprising comprising a combination of *Chlamydia pneumoniae* antigens. The combination of *Chlamydia pneumoniae* antigens may be one or more of the immunogenic compositions of the invention. The kit may further include a second component comprising one or more of the following: instructions, syringe or other delivery device, adjuvant, or pharmaceutically acceptable formulating solution. The invention also provides a delivery device prefilled with the immunogenic compositions of the invention.

EXAMPLES

The following invention will now be further described only by way of example in which reference is made to the following Figures. The following examples are presented only to illustrate the present invention and to assist one of ordinary skill in making and using the same. The examples are not intended in any way to otherwise limit the scope of the invention. Efforts have been made to ensure accuracy with respect to numbers used (e.g., amounts, temperatures, etc.), but some experimental error and deviation should, of course, be allowed for.

- Figure 1A. Assay of in vitro neutralization of C.pneumoniae infectivity for LLC-MK2 cells by polyclonal mouse antisera to recombinant Chlamydial proteins. Results are shown as reduction in the number of inclusions obtained when monolayers were infected with antiserum-treated infectious EBs, as compared to inclusion numbers given by untreated EBs. Percent reduction values are plotted against the reciprocal of the corresponding serum dilution. For each dilution inclusion counts were corrected for background inhibition of infectivity observed with the corresponding dilution of the pre-immune serum. The figure shows results obtained with serial dilutions of antibodies raised against a 'neutralizing' antigen (*), a 'non-neutralizing' FACS- positive antigen (v), and against the GST polypeptide, used in the fusion constructs, alone (σ).
 - Figure 1B shows serum titres giving 50% neutralization of infectivity for the 10 *C.pneumoniae* recombinant antigens described in the text. Each titer was assessed in 3 separate experiments (SEM values shown).
- Figure 2 shows immunoblot analysis of two dimensional electrophoretic maps of C.pneumoniae EBs using the imune sera described in the text. Immunoblots were obtained from either of two EB gels (panels A and B at the top) covering different pH intervals, according to which of the two allowed the best detection of a given antigen. The arrows in the HtrA immunoblot show which of the signals had a corresponding stained spot in the gel (arrows in panel A) which was subjected to MALDI-TOF identification. The two patterns in the HtrA blot are both suggestive of typical electrophoretic 'trains' composed of single charge variants of the same protein.
- Figure 3 shows mean numbers of *C.pneumoniae* IFU recovered from equivalent spleen samples from immunized and mock-immunized hamsters following a systemic challenge. Standard deviation values are shown above the bars. Antigens which induced significant protection are highlighted with an asterisk above the corresponding bar. All antigens were were delivered in Freund's adjuvant. n.i. = non immunized controls
- Figure 4 shows flow cytometric analysis of splenocytes from DNA-immunized HLA-A2 transgenic and non transgenic mice. Groups of 4 mice were immunized 3 times i.m. with 50μg of plasmid DNA expressing C. pneumoniae Low Calcium Response Protein H. IFN-γ production from splenocytes was monitored following either a 6h (ex-vivo) or a 6 day (restimulated) pulse with peptide CH-6 (10μg/ml). Equal numbers of gated live lymphocyte cells were acquired with a LSRII FACS System (Becton Dickinson) and percentages of IFN-γ producing CD8⁺ T cells were calculated using DIVA Software (Becton Dickinson).

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Figure 5 shows a flow cytometric analysis of splenocytes from transgenic and non transgenic mice infected with *C. pneumoniae* EBs. (A) HLA-A2 transgenic mice were intranasally infected twice with 5x10⁵ *C. pneumoniae* FB/96 EBs and splenocytes were stimulated for 6 days in the presence of relevant peptides before determining IFN-γ production by CD8⁺ T cells as described in the legend of Figure 4. (B) HLA-A2 transgenic and non transgenic mice were infected together with the same EBs preparation and CD8⁺ T cells were subjected to FACS analysis as reported in (A).

Table I shows a summary of data and properties of the C.pneumoniae antigens described in the text. The neutralization titer is reported is as the reciprocal of the antiserum dilution causing a 50% reduction in the number of inclusions in the in vitro infectivity assay. For the hamster model data the statistical significance of the results was evaluated by a two-tailed Student's t-test: significant data (p≤ 0.05) are highlighted with an asterisk. ND = not detected.

Table 2 shows results from hamster mouse model studies for hypothetical proteins.

Table 3 shows expressed genes of CPn EB selected by microarray.

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Table 4 shows C. pneumoniae selected peptides: protein sources and HLA-A2 stabilization assay.

Table 5 shows ELISPOT assay with CD8+ T cells from DNA immunised HLA-A2 transgenic mice.

Table 6 shows IFN- γ production from splenocytes of DNA immunized HLA-A2 transgenic and non transgenic mice.

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METHODS AND MATERIALS (Examples 1-4) (see Reference Section 1) C.pneumoniae EB purification

C.pneumoniae FB/96, a clinical isolate obtained from a patient with pneumonia at the Sant'Orsola Polyclinic, Bologna, Italy, was grown in LLC-MK2 cells seeded in individual wells of a six-well plastic plate (7). Cells were harvested 72 hr after infection with a sterile rubber, sonically disrupted and the elementary bodies (EB) purified by gradient centrifugation as described (26). Purified Chlamydiae were resuspended in sucrose-phosphate-glutamic acid (SPG) transport buffer, and stored in 0.5 ml aliquots, at -80°C until used. When required, prior to storage, EB infectivity was heat-inactivated by 3 hour incubation at 56°C.

Expression and purification of recombinant proteins

Open reading frames (ORFs), selected from the C. pneumoniae CWL029 genome sequence (16), were PCR-cloned into plasmid expression vectors and purified from E.coli cultures, essentially as previously described (25). Recombinant Chlamydial 15 proteins were obtained as GST fusion proteins by using pGEX-KG derived vectors (12) in E. coli BL21 (Novagen). PCR primers were designed so as to amplify genes without the N-terminal signal peptide coding sequence. When a signal peptide or processing site was not clearly predictable, the ORF sequence was cloned as 20 annotated by Kalman and coworkers (16). Recombinant E.coli cells were grown in LB medium (500 ml), containing 100 µg/ml Ampicillin, and grown at 37°C until $\mathrm{OD}_{600} = 0.5$, and then induced with 1 mM IPTG. Cells were collected by centrifugation 3 hr after induction and broken in a French Press (SLM Aminco, Rochester, NY). After centrifugation at 30.000 g, the supernatants were loaded onto Glutathione Sepharose 4B columns (Amersham Pharmacia Biotech) and column bound proteins were eluted with 50 mM Tris-HCl, 10 mM reduced glutathione, pH 8.0. Protein concentrations in the samples were determined using the Bradford method.

30 Preparation of mouse antisera

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Groups of four 5/6-week old CD1 female mice (Charles River, Como, Italy) were immunized intraperitoneally at day 1 with 20ug of protein in Complete Freund's adjuvant (CFA) and boosted at day 15 and 28 with 20ug of recombinant protein in Incomplete Freund's adjuvant (IFA). Pre-immune and immune sera were prepared from blood samples collected on days 0, 27 and 42. In order to reduce the amount of antibodies possibly elicited by contaminating E. coli antigens, the immune sera were incubated overnight at 4°C with nitrocellulose strips adsorbed with a total protein extract from E. coli BL21.

40 Flow cytometry assays

Analyses were performed essentially as previously described (25). Gradient purified, heat-inactivated EBs (2x105 cells) from C.pneumoniae FB/9, resuspended in phosphate-saline buffer (PBS), 0.1% bovine serum albumin (BSA), were incubated for 30 min. at 4°C with the specific mouse antisera (standard dilution 1:400). After centrifugation and washing with 200 µl of PBS-0.1% BSA, the samples were incubated for 30 minutes at 4°C with Goat Anti-Mouse IgG, F(ab)'2-specific, conjugated with R-Phycoerythrin (Jackson Immunoresearch Laboratories Inc.). The samples were washed with PBS-0.1%BSA, resuspended in 150 µl of PBS-0.1%BSA and analysed by Flow Cytometry using a FACSCalibur apparatus (Becton Dickinson, Mountain View, CA). Control samples were similarly prepared. Positive control

antibodies were: i), a commercial anti-C.pneumoniae specific monoclonal antibody (Argene Biosoft, Varilhes, France) and, ii), a mouse polyclonal serum prepared by immunizing mice with gradient purified C.pneumoniae EBs. Background control sera were obtained from mice immunized with the purified GST peptide used in the fusion constructs (GST-fusions control). FACS data were analysed using the Cell Quest Software (Becton Dickinson, Mountain View, CA). The shift between the background control histogram and the immune serum testing histogram was taken as a measure of antibody binding to the EB cell surface. The Kolmorov-Smirnov (K-S) two-sample test (44) was performed on the two overlapped histograms. The D/s(n) values (an index of dissimilarity between the two curves) are reported as "K-S score" in Table 1.

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2D Western Blot analysis of immune sera, and Mass Spectrometry Gradient purified C. pneumoniae EBs were washed with 5 mM Tris-HCl pH 7.5, 0.1 mM EDTA, 10% glycerol, centrifuged 15 min. at 13 000 x g and pellets were resuspended in reswelling solution (7 M urea, 2 M thiourea, 2% (w/v) CHAPS, (2%w/v) ASB14, 2% (v/v) IPG buffer pH 3–10 NL, or pH 4-7, 2 mM TBP, 65 mM DTT). Protein samples (200 or 20 μg of protein for Coomassie Blue stained reference gels, or gels to be processed for immunoblotting, respectively) were adsorbed overnight on Immobiline DryStrips (7 cm, pH 3-10 NL, or pH 4-7). Electrofocusing was performed in an IPGphor Isoelectric Focusing Unit (Amersham Biosciences, Uppsala, Sweden). The focused strips were equilibrated as described (15) and loaded on linear 9-16.5 % acrylamide gradients (7x 4 cm, 1.5 mm thick), for SDS-PAGE separation in a Mini Protean III Cell (Bio-Rad, Hercules, CA). Gels were stained with

colloidal Coomassie Blue (Novex, San Diego, CA) (4) and the protein maps so obtained were scanned with a Personal Densitometer SI (Molecular Dynamics) at 12 bits and 50 mm per pixel.

For Western Blot analyses, the proteins separated in the 2DE maps were transferred onto nitrocellulose membranes, overnight at 30 Volts, using a Protean III apparatus (BioRad, Hercules, CA). Membranes were stained with a 0.05% (w/v) CPTS (Copper(II) phthalocyanine-3,4',4",4"'-tetrasulfonic acid tetrasodium salt) in 12 mM HCl, and marked peripherally with 8 India-ink dots to provide anchors for subsequent image superimposition and matching. After scanning and image acquisition, the membranes were destained with 0,5 M NaHCO3, incubated with the mouse sera to be analyzed (either pre-immune or specific immune sera, diluted 1:1000), and then with a peroxidase-conjugated anti-mouse antibody (Amersham Biosciences, Uppsala, Sweden). After washing with PBS, 0.1% Tween-20, blots were developed using the Opti-4CN Substrate Kit (Biorad, Hercules, CA), and the images of the immunostained. blots again acquired as above. Images were analysed with the computer program Image Master 2D Elite, version 4.01 (Amersham Biosciences, Uppsala, Sweden). Superimposition and matches between Western-blot membranes and Coomassie stained gels were performed as follow. First, the CPTS-stained membrane image and the immunostained blot image were superimposed using the peripheral dot marks. Then, the sum image so obtained was superimposed to the Coomassie stained protein map using the CPTS stained CPn proteins as anchors. The areas on the Coommassie stained map corresponding to immunostained spots on the blot were excised from the preparative gel for protein identification. Protein sample were dried in a vacuum centrifuge, and in-gel digested, for 2h at 37°C, with an excess of porcine Trypsin (Promega, Madison, WI), in 100 mM ammonium bicarbonate. Tryptic peptides were desalted and concentrated using Zip-Tip (Millipore, Bedford, MA). Peptides were

directly eluted and loaded onto a SCOUT 384 Anchor Chip multiprobe plate (400 µm, Bruker Daltonics, Bremen, Germany) with a solution of 2-5 dihydroxybenzoic acid (5g/l), in 50% acetonitrile, 0.1% trifluoroacetic acid. Spectra were acquired on a Bruker Biflex III matrix-assisted laser desorption ionization-time of flight (MALDITOF) apparatus. Resulting values for monoisotopic peaks were used for database searches using the Mascot software (32), as available at the website http://www.matrixscience.com/.

In vitro neutralization assays

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10 In vitro neutralization assays were performed on LLC-MK2 (Rhesus monkey kidney) epithelial cell cultures. Serial four-fold dilutions of mouse immune and corresponding preimmune sera were prepared in sucrose-phosphate-glutamic acid buffer (SPG). Mouse polyclonal sera to whole EBs were used as positive control of neutralization, whereas SPG buffer alone was used as negative control of neutralization (control of 15 infection). Purified infectious EBs from the C.pneumoniae FB/96 were diluted in SPG buffer to contain 2.5x10⁷ IFU/ml, and 10ul of EBs suspension were added to each serum dilution in a final volume of 100ul. Antibody-EB interaction was allowed to proceed for 30 min at 37°C on a slowly rocking platform. The 100ul of reaction mix of each sample was used to inoculate PBS-washed LLC-MK2 confluent 20 monolayers (in triplicate for each serum dilution), in a 24-well tissue culture plate, and centrifuged at 805 x g for 1 hour at 37°C. After centrifugation Eagle's minimal essential medium containing Earle's salts, 20% fetal bovine serum and lug/ml cycloheximide was added. Infected cultures were incubated at 37°C in 5%CO₂ for 72 hours. The monolayers were fixed with methanol and the Chlamydial inclusions were 25 detected by staining with mouse anti-Chlamydia fluorescein-conjugated monoclonal antibody (Merifluor Chlamydia, Meridian Diagnostics, Inc.) and quantified by counting 10 fields per well at a magnification of 40X. The inhibition of infectivity due to EBs interaction with the immune sera was calculated as percentage reduction in mean IFU number as compared to the SPG (buffer only)/EBs control. In this 30 calculation the IFU counts obtained with immune sera were corrected for background inhibition of infection due to the corresponding pre-immune mouse serum. According to common practice, the sera were considered as "neutralizing" if they could cause a 50% or greater reduction in infectivity. The corresponding neutralizing titer was defined as the serum dilution at which a 50% reduction of infectivity was observed. Experimental variability was evaluated by calculating the standard error of 35 measurement (SEM), from three titration experiments for each recombinant antigen, as shown in Fig.1B.

In vivo screening

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In vivo evaluation was performed using a hamster model of systemic infection, as recently described (34). Essentially, adult (10-11 week old) Syrian hamsters (Morini, S. Polo D'Enza, Italy), previously immunized with the recombinant vaccine candidates were challenged systemically with infectious Cpn elementary bodies (EB). Protection was assessed by determining the levels of viable EB recovered from the spleen, as compared to non-immunized animals. Statistical significance of the results was evaluated by a two-tailed Student's t-test.

Groups of 8 hamsters were immunized subcutaneously with recombinant antigens, or only injected with buffer for the control group, at days 0, 7, and 21. For each immunization 20 ug protein 1:1 diluted with Freund's complete adjuvant (first dose)

and Freund's incomplete adjuvant (booster doses) was injected. At day 35 post-infection the hamsters were anaesthetised with Ketamine and inoculated intraperitoneally and intranasally with 0.1 ml of *C.pneumoniae* EB suspension (1.0x10⁸) at each site. Animals were sacrificed seven days after infection. The spleen was weighed, and homogenized in a mortar to obtain a 10% (wt/vol) suspension in cold SPG buffer. Tissue suspensions were centrifuged at 300 x g for 10 min at 4°C to remove coarse debris. The clarified homogenates (0.2 ml) were inoculated in duplicate onto LLC-MK2 cells seeded in plastic individual well of a 24 well plate, incubated at 37°C for 72 h and fixed in acetone before detection and counting of numbers of *Chlamydia*l inclusions per well by immunofluorescence microscopy. The protocol was approved by the ethical committee of the University of Bologna.

Example 1 (in vitro studies)

Screening antisera for in vitro neutralizing properties

Following a genome-wide screening for proteins likely to be localized on the cell surface of C. pneumoniae, we recently reported (25) that antisera to 53 recombinant Chlamydial antigens were capable to bind in a FACS assay, the surface of Chlamydial cells. In order to check whether some of the FACS positive antigens were capable of interfering with EB in vitro infectivity, we raised mouse antisera against the recombinant FACS positive antigens and assessed the effect of each antiserum on the infectivity of purified EBs with respect to monolayers of LLC-MK2 cells. Infectious EB were first incubated with the antiserum and then used to infect cell monolayers in 24-well multititer plates. In parallel, control samples were similarly processed in which the EBs were: i), either treated with buffer only, or, ii), treated with the same dilutions of the corresponding preimmune mouse sera.

15 Results I

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Using this assay, 10 sera have so far proved to effectively neutralize in vitro infectivity to an extent greater than 50%, a property that common practice qualifies such antigens as "neutralising" (Figure 1). These 10 sera were obtained by mouse immunization with recombinant proteins derived from the following *C.pneumoniae* genes:

- pmp10 and pmp2, encoding two members of the heterogeneous Chlamydial PMP family of polymorphic membrane proteins;
- artJ, encoding a putative extracellular solute (possibly Arginine) binding protein of an aminoacid transport system;
 - eno, encoding a protein homologous to bacterial enolases, glycolytic enzymes which can be found also on the bacterial surface:
 - htrA, encoding a putative chaperone with heat-shock inducible protease activity;
- the Cpn0301 "hypothetical" gene, encoding a protein homologous to the ompH family of bacterial proteins, some members of which have been shown to be chaperones involved in outer membrane biosynthesis:
 - two Cpn-specific "hypothetical" genes Cpn0795 and Cpn0042;
 - omcA encoding a 7-9 kDa protein annotated as an outer membrane protein; and
 - atoS a putative sensor member of a transport system.

As shown in Figure 1 and summarized in Table I, OmpH, enolase and Cpn0795 appeared to induce the highest neutralizing sera, with titers above 400. By contrast, Pmp2, ArtJ and Cpn0042 induced titers equal or less than 100, while the remaining 4 antigens, Pmp10, HtrA, AtoS and OmcA showed intermediate titers.

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Example 2 (in vivo studies)

Evaluation of antisera specificity by 2D immunoblot analysis of Cpn protein extracts

- In order to investigate if the neutralizing activity observed in the *in vitro* infection of LLC-MK2 monolayers was actually due to the binding of the antibodies to the selected *C.pneumoniae* proteins, rather than to possible cross-rections with other antigens, we assessed the specificity of the antisera by immunoblot analysis of two dimensional electrophoretic maps of EB proteins.
- In particular, this analysis was carried out on six antigens (Pmp2, Pmp10, Eno, ArtJ, 10 HtrA and OmpH-like) known to be visible in the 2D maps of EB total proteins (Montigiani et al., 2002 Infection and Immunity 70: 368-379). Total EB proteins were resolved by 2D-electrophoresis using two different pH intervals (pH 3-10 non linear, and pH 4-7, respectively) since previous experiments had shown that some of the proteins under study were better detected using one rather that the other of the 15 above pH intervals. For each pH interval four gels were run in parallel. One gel was stained with Coomassie Blue to visualize the protein spots, while the other gels were blotted on nitrocellulose filters and stained with one of the selected sera at 1000-fold dilution. Subsequently, the images of the immunostained blots (Fig.2, panels c to h) 20 were superimposed to the corresponding Coomassie Blue-stained gel to identify the spots which had reacted with a given antiserum. The matching protein spots were excised and processed for peptide identification by MALDI-TOF analysis.

Results 2

- In all six maps the immunoreactive protein species in the excised gel area were found to contain peptides from the expected *Chlamydial* protein. Even when the serum reacted with more than one electrophoretic protein species, the mass spectra of all spots which could be detected in the Coomassie Blue stains 2DE map were always consistent with the same polypeptide being present as multiple electrophoretic species.
- Interestingly, the immunoblot obtained with the HtrA antiserum showed two sets of 4 spots arranged as two typical electrophoretic 'trains' at two different molecular weights. On the Coomassie Blue stained gel it was possible to identify 4 corresponding spots, 3 in the upper train and 1 in the lower Mw set. MS analysis identified all of them as products of Cpn HtrA gene. Interestingly the lower Mwt species missed 3 N-terminal tryptic peptides, detected in the higher Mw spot series, and mapping within the first 100 aa of the ORF. These results suggest that HtrA was present in the EB protein sample both as a full htrA product, and as a discrete species missing a short N-terminal peptide, possibly as a result of some post-translational processing.

Discussion of Results 2

In the analysis of data which are based on polyclonal antibody reactivity one should consider that cross-reactions due to epitope mimickry are always difficult to exclude.

The problem of antisera specificity was addressed in this work by 2D immunoblotting and identification of the reacting electrophoretic species by mass spectrometry analysis. This approach was possible for 6 of the 10 antisera, i.e. those corresponding to proteins previously identified by mass spectrometry (MALDI-TOF) analysis on 2D electrophoretic maps of *C.pneumoniae* EB proteins (25, 42) (Table 1, and Figure 2).

The probability of fortuitous cross-reactions between unrelated *Chlamydial* protein

species was minimized by the results of the immunoblotting analyses which showed that out of ca 300 protein spots in a map, all those reacting with the tested antisera were consistent with the expected antiserum specificity. Obviously, since during 2-D electrophoresis conformational epitopes are generally lost, structure-dependent cross-reactions cannot be ruled out in this type of analysis.

Example 3

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In vivo evaluation of the in vitro neutralizing antigens in a hamster model of systemic infection

We have recently described a new hamster model of systemic Chlamydia pneumoniae infection in which replicating Chlamydia disseminate through macrophages and accumulate in the spleen (34). We therefore asked the question whether the in vitro neutralizing antigens we identified would also have protective activity in vivo using this model. To this aim, the 10 in vitro neutralizing recombinant antigens were used to immunize 8 hamsters with 3 subcutaneous injections over a three-week period, and challenged with 2x10⁸ Cpn EBs two weeks later. Spleen infection was assessed 7 days after challenge. The difference between the mean number of infectious Chlamydiae recovered from control animals and the mean number of Chlamydiae recovered from animals immunized with the recombinant Chlamydial antigens, was taken as a measure of protection specifically induced by the putative vaccine candidate.

Results 3

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The results of spleen protection observed for the various antigens in repeated experiments are shown in Figure 3 and reported as percentage values in Table 1. Six out of ten antigens, Pmp2, Pmp10, Enolase, the OmpH-like protein, and the products of the *C.pneumoniae*-specific genes Cpn0759 and Cpn0042, showed a statistically significant protective activity, with a reduction in IFU recovered from the spleens of immunized animals higher than 80% with respect to mock-immunized controls.

A limit of the hamster model is that, because of the absence of immunological reagents, the relative contribution of humoral and cell-mediated immunity cannot be assessed. However, we asked the question whether recombinant antigens could elicit also in the hamster neutralizing antibodies with sufficiently high titers. Therefore we tested the sera from hamsters immunized with Pmp2 and enolase, two of the most protective antigens, in the *in vitro* neutralization assay. Both antigens had a neutralizing titer of approximately 100 (data not shown).

Summary of Results 3

In conclusion, a considerable proportion (60%) of the *in vitro* neutralizing antigens were also protective in the hamster *in vivo* model and our data suggest that antibody-mediated neutralization could play a role at least in this model of systemic infection.

Discussion of Results 3

Beside assaying the *in vitro* neutralization properties of the selected subset of 10 FACS-positive antigens, we also assessed the performance of these antigens in protecting against *C. pneumoniae* infection in an animal model of systemic infection recently described in the hamster (34). This evaluation addressed the capability that the recombinant antigens would have of inducing a protective response against naturally replicating *Chlamydiae* (rather than EB's purified from *in vitro* cultures grown under artificial conditions) and in the context of a systemic infection. In fact

the hamster model we used, while it does not model the typical respiratory infection considered to be the predominant route by which *C. pneumoniae* infects humans, it nevertheless simulates a situation of systemic invasion which could be preliminary to the establishment of *C. pneumoniae* chronic infection considered by several authors as being associated to the development or progression of cardiovascular disease, and other chronic degenerative diseases. Notably, a limit of any hamster model is the current lack of hamster-specific immunological reagents which would allow the analysis of cell mediated immune responses. However, in the case of systemic infections, by common wisdom, neutralizing antibodies are likely to have a protective action. The finding that 6 of the 10 'in vitro neutralizing' antigens had also a >80% protective action in vivo, and that a measurable neutralizing activity was also found in the sera of immunized hamsters, suggests that a specific antibody mediated immunity could at least partially contribute to the animal protection here described.

15 Example 4

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Two 'hypothetical' proteins 6784 and 6814 (encoded by the ORFs Cpn0498 and Cpn0525) yielded FACS-positive sera which, however, were not able to neutralize host cell infection *in vitro*. However, these antigens performed remarkably well in the hamster-spleen test.

Table 2

Gene/ORF ID in CWL029	Protein ID	Recombina nt Fusion Type	Annotation	Recipr ocal of 50% neutral isation titre	% Protection in the hamster spleen test (ref 34)
Cpn0498	4376784	GST	Hypothetical protein	0	94
CPn0525	4376814	GST	Hypothetical protein (similarity to CT398)	0	97
CPn0324		HIS	Low Calcium Response Element (LcrE)	·	Completely protected 8 of 16 animals and reduced the infectivity titres of the eight positive animals

Discussion of Results 4

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Interestingly, whilst antiserum against CPn0525 gave negative in vitro results (ie no neutralising activity), the CPn0525 protein gave 97 per cent protection from spleen infection in an in vivo hamster immunisation assay (ie a positive in vivo result). Likewise, whilst antiserum against Cpn0498 gave negative in vitro results (ie no neutralising activity), the CPn0498 protein gave 94 per cent protection from spleen infection in an in vivo hamster immunisation assay. Thus a positive signal obtained in the FACS assay does not guarantee a corresponding positive in vitro neutralization activity and conversely a negative neutralization activity does not mean that a positive in vivo result can be excluded.

General Discussion of Results 1-4

Strategy for identification of Chlamydia pneumoniae antigens of interest
Our strategy was based on the following experimental steps: 1) analysis of
Chlamydia genome sequence to select putative membrane-associated antigens, 2)
cloning, expression and purification selected antigens, 3) preparation of antigenspecific sera by mouse immunization with the purified antigens, 4) FACS analysis of

Chlamydia EBs using the mouse sera to identified surface-exposed antigens, 5) "in vitro neutralization" assay to test whether antibodies elicited by a given antigen can interfere with the process of eukaryotic cell infection, and 6) use of appropriate animal model to test the capacity of selected antigens to confer protection against Chlamydia challenge.

As recently described by Montigiani et al ((2002) Infection and Immunity 70: 368-379) from the initial screening of the C.pneumoniae genome, a panel of mouse sera 25 was prepared against over 170 recombinant His-tagged or GST-fusion proteins encoded by genes or "open reading frames" somehow predicted to be periferally located in the Chlamydial cell. When these antibodies were tested in a FACS assay for their ability to bind the surface of purified C.pneumoniae EBs, a list of 53 "FACS-30 positive" sera was obtained. The corresponding putative surface antigens were then further assessed for their capability of inducing neutralizing antibodies. This part of the work involved testing which of the sera contained antibodies capable of interfering with the process of in vitro infection of epithelial cell cultures. In the in vitro "neutralization" assay purified infectious EBs are incubated with progressive 35 dilutions of the immune sera and, in parallel, dilutions of the corresponding preimmune sera, and of sera against non Chlamydia control antigens.

Cell cultures are infected in the presence of cycloheximide, which inhibits host cell protein synthesis and favours *Chlamydia*l intracellular growth with the consequent formation of typical cytoplasmic inclusions which can be stained with *Chlamydia* specific fluorescence-labeled monoclonal antibodies and counted with an UV light microscope. Working with appropriate pathogen-to-host cell ratios, it can be reasonably assumed that the number of detected cytoplasmic inclusion is proportional to the number of infectious *Chlamydiae* in the original sample. So a reduction in inclusion numbers caused by the presence of an antigen-specific antiserum, as compared to the numbers obtained with control sera, gives a measure of the capability of a given antigen to elicit antibodies which can inhibit some stage of the *Chlamydia*l infection process. According to common convention, an anti-serum is labelled as 'neutralizing' when the reduction of infectivity is equal or greater than 50%, and the

serum dilution yielding a 50% reduction in infectivity is referred to as the 50% end-point neutralization titer.

Some of the results obtained by screening the panel of recombinant antigens with the C.pneumoniae in vitro neutralization assay confirm that some of the listed antigens, 5 like the members of the family of heterogeneous polymorphic membrane proteins (PMP), which, on the basis of published literature data, could be reasonably expected to be surface-exposed and possibly induce neutralizing antibodies. However, there are also proteins which could be considered so far only hypothetical, and proteins which just on the basis of their current functional annotation could not be at all 10 expected to be found on the bacterial surface. Using an in vitro neutralising assay, it was found that sera to 10 CPn antigens have so far proved to effectively neutralize in vitro infectivity to an extent greater than 50%, a property that common practice qualifies such antigens as "neutralising" (Figure 1). These 10 sera were obtained by mouse immunization with recombinant proteins derived from the C.pneumoniae 15 genes listed below.

Using a recently described *in vivo* model of systemic infection (hamster model), hamsters immunised with 6 of the *in vitro* neutralising antigens, when challenged with CPn EBs, showed a greater than 80% reduction of spleen infection as compared with non-immunised controls.

Characterisation of 10 CPn proteins

The proteins identified by the present work can be divided in 3 groups:

- proteins which have an annotation compatible with (could be reasonably expected to have) an expected/predicted exposure on the *Chlamydial* cell surface and with the possibility that antibodies binding to them may actually interfere with host cell attachment and entry (ie proteins which could possibly induce neutralising antibodies)
- proteins which by homology with other gram-negative bacteria could be expected to have a periplasmic exposure (ie would not be expected at all to be found on the bacterial cell surface); and
 - proteins which are still labelled as 'hypothetical' (ie cellular location and/or cellular function not known)

35 *Group 1*

(Pmp proteins (pmp2 and pmp10), OmcA and OmpH)

Pmp proteins (pmp2 and pmp10)

The first group includes the 2 polymorphic outer membrane proteins (Pmp's) Pmp2 and Pmp10 (10, 11, 14, 30), the outer membrane protein OmpH-like, and OmcA, 40 annotated (Chlamydia Genome Project at http://Chlamydiawww.berkeley.edu:4231/) as "predicted 9-kD cysteine-rich, outer membrane protein, lipoprotein". The Pmp family of Chlamydia-specific proteins is generally thought to comprise probable pathogenicity factors, with an autonomous secretion capacity (autotransporters), important for adhesion to host cells and are generally considered as 45 promising vaccine candidates. However, apart from very recent unpublished results on Pmp21, this is the first time that antisera to recombinant Pmp's are reported to have neutralizing properties.

OmcA.

OmcA is the product of a gene co-transcribed in the same operon with the 60 kDa OmcB cystein-rich protein which is a major structural component of the *Chlamydial* outer membrane and a major immunogen in human *C. trachomatis* infections. OmcB and OmcA are likely to interact in some as yet unknown outer membrane structure, so it is possible that antibodies to OmcA can interfere with EB infectivity.

OmpH

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Finally, the Chlamydial OmpH is probably a member of the OmpH (Skp) family of proteins which have been reported to have chaperonin activities in other bacteria very important for the correct biosynthesis of the outer membrane. These proteins appear to cooperate in this task with HtrA (see below). In fact, in E.coli single KO mutants of either OmpH (Skp) or HtrA (DegP) are still viable, but double mutants do not grow (37). It should be pointed out that even if the amino acid sequences of the ompH-like proteins of Chlamydia (all C.pneumoniae and C.trachomatis or C.caviae variants) line-up very well with the rest of the bacterial OmpH proteins, they are the only ones to be acidic, whereas the rest of the family comprises mostly very basic proteins (including some with histone like behaviour, at least in vitro). One could speculate that if the chaperone activity is maintained also in the ompH like Chlamydial proteins, this may be related to some Chlamydial peculiarity.

Second Group of Selected Proteins (ArtJ, AtoS, HtrA and Enolase)

The second group, which represents a somehow surprising finding, includes ArtJ, AtoS, HtrA and Enolase. If the current annotation (justified by analogy with homologous genes in other bacteria) is correct, all these proteins would be expected to have a periplasmic location in gram-negative bacteria. and to be surface-exposed only in a gram-positive bacterium. It is possible that owing to their atypical life cycle, requiring an efficient passage from a dormant spore-like status (the EB) to an active form needing to adapt quickly to host-cell responses to invasion, *Chlamydiae* in fact display some sensors directly on the outer surface of their infectious form.

ArtJ

In the case ArtJ – for which we have data supporting both antigen expression and serum specificity – the hypothesis of an atypical situation peculiar to *Chlamydia* is supported by the anomalous gene set-up resulting from the present analysis of the *Chlamydia* genomes. ArtJ is so annotated by analogy with the ART transport systems of *E.coli* which has 5 genes organized in two operons (24): artPIQM and artJ which are responsible for the arginine transport. In Cpn however the artPIQM genes are absent and therefore it appears that *Chlamydia*l ArtJ operates in a molecular context which is different from its *E.coli* model and must be peculiar to this species.

HtrA

HtrA (DegP), which in other bacteria has a complex hexameric structure, has been described as having multiple functions (3, 5, 18, 19, 27, 38): a chaperonin assisting a correct outer membrane biogenesis, inducible protease for the elimination of misfolded membrane proteins, and also a sensor of 'stress' conditions. In *Chlamydia* none of these properties has been demonstrated yet, however we find that in purified EB HtrA is present in two forms one of which appears to be processed by being deprived of the N-terminal fragment. This fragment, if aligned with the homologous

HtrA sequence from Thermologa maritima (18), would comprise a predicted loop acting as a structural lid controlling the access to the protease active. So it appears tempting to speculate that HtrA could have a similar protease activity and the two forms identified on the 2-D map represent the active and inactive species. Interestingly, the *C. trachomatis* HtrA ortholog is recognized by human sera from patients who had a *Chlamydial* genital infection (35), and a similarly HtrA is one of the antigens in the immunoproteome of Helicobacter pylori (13). Furthermore the homologue protein in *Haemophilus influenzae* is a protective antigen in both a passive infant rat model of bacteremia and the active chinchilla model of otitis media (23).

Enolase

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Also in the second group of proteins expected to be located elsewhere than the cell surface, is Cpn enolase. This protein aligns with the well known family of conserved glycosylases, which are essentially cytoplasmic enzymes, but in Streptococci enolase has been shown to have also a cell surface location, and extracellular matrix binding properties (1, 28, 29)). Interestingly, Gaston and colleagues (8) also showed that in patients with reactive arthritis induced by *C.trachomatis*, enolase induces specific CD4⁺ T-cell responses. Furthermore, a clone responding to the enolase *C.trachomatis* ortholog, responded also to *C.pneumoniae* EBs, and, since no proliferative response could be observed using a fungal or a mammalian enolase, the authors of this study concluded that the CD4 T-cell stimulating epitope must be *Chlamydia* specific.

Third Group of Proteins

(unknown cellular location and/or cellular function, Cpn0759, CPn0042)

25 The third of the 3 groups in which we propose to divide, just for the sake of discussion, the 10 neutralizing antigens above described, comprises two proteins which are still annotated in public Chlamydial databases as the hypothetical products of two CPn-specific genes: Cpn0759 and Cpn0042. The Cpn0759 gene is the second gene in a cluster of 6 Cpn-specific hypothetical genes (from Cpn0794 to Cpn0799) 30 immediately upstream of the enolase gene. With the exception of Cpn0759 the products of all the other genes in the cluster share similarities of 30 to 40% over long stretches of amino acids. The Cpn0042 gene encodes a hypothetical protein, with 4 coiled-coil regions, which has been described as a member of a new family of hypervariable outer membrane proteins (33). Interestingly, the hypervariability of 35 these proteins could be due to a strand-slippage mechanism induced by the presence of a poly(C) stretch within the coding region of the corresponding genes, a mechanism already described in the Pmp's family for the pmp10 gene (30). However, as indicated by their annotation, the function of these proteins is still unknown, and our observations provide the first experimental indication of a possible function 40 related to the Chlamydial infection process.

Fourth Group of Proteins Cpn0498

So in this case the triple parallel-screening evaluation, with two positive and one negative result, brought to the identification of a previously unknown antigen (ie an antigen with unknown biological function) having, according to current views, just the desirable basic properties in terms of antigenic function of a vaccine candidate.

Example 5

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Background

The main stages in the *Chlamydial* life cycle are:

- (i) the binding to the host cell surface and entry into the cytoplasm through a specialised vacuole (the *Chlamydial* inclusion) by an extracellular spore-like infective form, called the elementary body (EB); and
- (ii) the conversion of the EB to a non-infective replicative form called a reticulate body (RB) that replicates by binary fission a number of times within the inclusion to form a microcolony.
- 10 The sets of genes which are expressed in the various stages of the *Chlamydial* life cycle and the signals that trigger the passage from one stage to the next are largely unknown and still need investigation.
- Current DNA microarray techniques permit profiling of gene expression at the mRNA level as a function of the cellular state. This can lead to the identification of genes or 15 clusters of genes whose up- or down-regulation is associated to a particular state of the cell and to the identification of therapeutically relevant targets. Using this technology, DNA fragments representing specific portions of all genes belonging to a given organism (the fragments can be derived from cDNA libraries or can be obtained 20 by PCR amplification and chemical synthesis) are chemically bound to the surfaces of solid supports (chips) at high densities and in an ordered manner. Currently up to 10, 000 DNA fragments or 250, 000 oligonucleotides can be spotted onto a single squared centimetre of chip surface. The DNA chips are then utilised to define which of the spotted genes are transcriptionally active in a particular cellular population. To do so, 25 RNA is prepared, labelled with fluorescent dyes and finally hybridised to the DNA fragments fixed to the surface of the chip. By using an appropriate computer-assisted fluorescence detector, the fluorescence signals emitted by each spot upon excitation with a laser beam is carefully quantified to define the transcription activity of all the arrayed genes.

CPn DNA microarrays have been developed to look at the transcriptional events which occur when a given CPn pathogen gets into contact with the host cells, both in *in vivo* and *in vitro* settings. DNA chips carrying the entire genome of a particular bacterium, such as the CPn bacterium can be prepared in a very short period of time so that whole genome expression analysis can be determined.

Experimental Methodology

Specifically, a genomic DNA (open reading frame probes) microarray approach for gene expression in CPn bacteria was adopted. In this respect, an array was prepared for the analysis of the CPn life cycle on the basis of the published annotation of the complete genome. The *Chlamydia* DNA chips carry about 1000 PCR-derived DNA fragments, which have an average size of 400-700bp and correspond to internal portions of all CPn annotated genes.

45 Results 5

Table 2 shows transcriptional activity for expressed genes for CPn EB selected by microarray. The data in Table 2 shows different mRNAs in order of abundance from cells in their infectious "spore-like" (EB) form The cells were used at the end of their cycle where gene expression is likely to be at its highest. As values less than 300 is

likely to be background, the top set of proteins (approx top 30) with more intense signals are likely to be the most interesting proteins
A review of the values for the expressed genes indicates that three of the FACS positive CPn antigens (CPn0331 (hypothetical), CPn0234 (hypothetical) and CPn0572 (hypothetical) are all highly expressed genes.

Materials and Methods (Examples 6-8) (Reference Section II) T cell Epitope prediction and peptide synthesis

- T cell epitope prediction was carried out on the genomic sequence of *C. pneumoniae* CWL029 strain (Accession numbers NC 000922 or AE001363) using the BIMAS algorithm [24]. Synthetic peptides (purity > 80%) were synthesized by Primm Srl (Milan, Italy), suspended in 100% DMSO and kept at -20° C before use.
- 15 RMA-S/A2 cell line and HLA-A2 transgenic and non transgenic mice

 The T cell lymphoma murine cell line RMA-S stably transfected with HLA-A2 (RMA-S/A2, H-2^b, TAP2), was kindly provided by Dr. Barnaba, Università degli Studi "La Sapienza", Rome, Italy, and cultured at 37° C in RPMI-1640 (GIBCO) supplemented with heat inactivated 10% FCS, 100 IU/ml penicillin/streptomycin, 2

 20 mM Lglutamine (GIBCO) and 5×10-5 M 2-ME (Sigma). H2-b HLA-A2 transgenic mice [35] were housed in a pathogen-free environment and screened for HLA-A2 expression by FCM carried out on total blood samples using the BB7.2 anti-A2 mAb [48]. Only mice with percentages of A2 expressing cells higher than 70-80 % were used for DNA immunization and C. pneumoniae infection experiments. Animals which showed no HLA-A2 expression were mated in order to obtain an HLA-A2 non transgenic population, to be used as a control in the experiments.

Epitope stabilization assay

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RMA-S/A2 cells (3-5 x 10⁵/well) were seeded in serum-free RPMI medium, supplemented with human β2 microglobulin (3 μg/ml, Sigma), without or with the test peptide (10μM). Following overnight incubation at 26°C in humidified 5% CO₂ atmosphere, cells were shifted to 37° C for 2 h before determining the HLA-A2 expression level at the cell surface using the BB7.2 anti-A2 mAb and a PE-conjugated anti-mouse IgG (Jackson ImmunoResearch). Fluorescence intensity on living cells, which did not incorporate propidium iodide, was analyzed by FCM. As controls, corresponding samples without peptide and samples with peptide but treated only with the anti-mouse secondary antibody, were used.

Infection and DNA immunization of HLA-A2 transgenic and non transgenic mice

Transgenic mice were intranasally infected twice with a month interval, using 5×10^5 C. pneumoniae FB/96 EBs [4] diluted in 50 µl of PBS. C. pneumoniae antigen coding genes were amplified by PCR using FB/96 genomic DNA, cloned into plasmid pcmvKaSF2120 [49] and verified by DNA sequence analysis. Fifty µg of endotoxin free recombinant plasmid DNA, diluted in Dulbecco's phosphate buffer (GIBCO), were injected into the tibialis muscle of mice at days 0, 21 and 35.

CD8⁺ T cells isolation and IFN-γ determination by ELISpot assay
Splenocytes from DNA immunized mice were prepared one week after the third
immunization using Cell Strainer (Falcon) filters. Following red blood cells lysis,

CD8⁺ T cells from spleen cells suspensions were enriched by positive selection using magnetic activated cell sorting (MACS-Miltenyi Biotec) with CD8a (Ly-2) microbeads. CD8⁺ T cells purity was higher than 90%, as determined by FMC. Multiscreen 96-well nitrocellulose plates (Millipore) were coated with 5 µg/ml of the anti-mouse IFN-y antibody (R4-6A2, PharMingen) in 100 µl of carbonate buffer, pH 9.2. After overnight incubation at 4°C, plates were saturated at 37°C with 200 ul of BSA (1%) in PBS for 2 h. Purified CD8⁺ (5x10⁴) were added in a total volume of 200 µl/well in the presence of irradiated (3,000 rad) spleen cells from non immunized HLA-A2 transgenic mice as a source of antigen-presenting cells (2x10⁵/well), 10 10 μg/ml of peptide and 10U/ml of human r-IL-2 (Chiron Corporation). After incubation for 20 h at 37° C, 5% CO₂, plates were washed and developed for bound IFN-γ using sequential incubations with biotinylated antimouse IFN-γ (XMB1.2, PharMingen), ExtrAvidin-alkaline phosphatase and substrate BCIP/NBT (Sigma) dissolved in water. Spots were enumerated in each well using a dissecting microscope. Medium 15 containing an irrelevant peptide or without peptide was used as negative controls, while positive controls were represented by CD8⁺ T cells treated with ConA (5 μg/ml).

In vitro cultures and flow cytometric analysis of splenocytes from transgenic and non transgenic mice infected with C. pneumoniae

Splenocytes from infected mice were isolated one week after the second infection with C. pneumoniae Ebs. For ex vivo analysis of IFN-y production, 2x10⁶ splenocytes were seeded in the presence of the test peptide (10µg/ml) and anti-mouse CD28 antibody (1µg/ml, PharMingen) as co-stimulus. After a two h incubation at 37° C, 5 % CO₂, Brefeldin A (10 μg/ml, Sigma) was added and the incubation was extended for 4 additional hours. Following two washes with PBS, cells were permeabilized, fixed and stained both with anti-murine-IFN-γ-(PE), anti-murine CD8 (APC) and antimurine-CD69 (FITC) and the corresponding isotypes. Positive controls for cytokine production were performed on cells stimulated with anti-mouse CD3 and CD28 antibodies (1 µg/ml, PharMingen). Cells cultured either in the absence of peptide or pulsed with the HepB negative control peptide were used as negative controls. All samples were analyzed using a FACS LSRII flow cytometer (Becton Dickinson). For analysis of IFN-γ production by short term T cell lines, 5-10x10⁶ splenocytes from infected mice were cultured for 6 days in the presence of the test peptide (20 µg/ml), with rIL-2 (10 µg/ml) being added after the first two days. At the end of the incubation period, cells were washed twice in RPMI, pulsed again for 6 h in the presence of the test peptide (10µg/ml), 1x10⁵ freshly prepared CD8 depleted antigen presenting cells from HLA-A2 transgenic mice (irradiated at 3000 rad) and antimouse CD28 antibody (1µg/ml, PharMingen) as co-stimulus. After a two h incubation at 37° C, 5 % CO2, Brefeldin A (10 µg/ml, Sigma) was added, the incubation was extended for 4 additional hours and IFN-y production was analyzed by FCM.

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Example 6

In silico analysis of *Chlamydia pneumoniae* genome and prediction of HLA-A2 T cell epitopes

The genome of the Chlamydia pneumoniae CWL029 strain was used to predict 9mer peptide sequences with high probability to bind class I HLA-A2 molecules. The analysis was carried out using the predictive algorithm available at the NIH Bioinformatics & Molecular Analysis Section Web server (http://bimas.cit.nih.gov/), which ranks potential MHC binders according to the predictive half-time dissociation of peptide/MHC complexes [24]. Since some Chlamydial proteins have been reported 10 to induce autoimmune responses [25-28], we restricted our search to a subset of proteins, distinctive of the Chlamydia genus, consisting of 13 protein identified as members of the type III secretion system, 17 Polymorphic Membrane Proteins (PMP) and 19 additional proteins, 5 of which already identified as EB surface antigens [4]. The predicted binding score of 157.22, obtained for the well characterized HIV-1 p17 gag epitope ⁷⁷SLYNTVATL⁸⁵ [29], was taken as an arbitrary cut-off for peptide 15 selection. A total of 55 potential C. pneumoniae-derived T cell epitopes, belonging to 31 different proteins, were identified (Table I), which had predicted binding scores ranging from 156.77 to 42,485.263

20 In vitro binding of peptides to HLA-A2

The capacity of the selected peptides to bind to HLA-A2 was assessed using an in vitro MHC class I stabilization assay, carried out with the murine transporter associated with antigen processing (TAP)-deficient cell line RMA-S/A2, stably transfected with the human class I A2 gene. MHC class I molecules, cultured at 37° C, are unstably expressed on the cell surface of TAP-deficient cells [30-32]. Culturing 25 the cells at 37° C in the presence of binding peptides, results in formation of a more stable MHC/peptide complex which can be monitored by flow cytometric analysis. RMA-S/A2 cells were therefore cultured overnight at 26° C in the presence of the test peptides, shifted to 37° C for 2 hours and the surface level of stabilized A2 30 molecules was quantified by direct staining with an anti-HLA-A2 specific mAb. Two known HLA-A2 restricted CTL epitopes were used as positive controls for binding to A2, the HIV-1 p17 gag peptide [29] and the influenza matrix M1 protein peptide FluMP58 [33], while the Hepatitis B virus envelope antigen peptide HbenvAg125 (HepB) was used as a negative control [34]. 35

Results 6

The binding results obtained are shown in Table 4 and allowed the identification of 15 peptides with a net mean fluorescence intensity (Net MFI) higher than 92.3, corresponding to the value obtained with the HIV-1 p17 gag positive control peptide, 8 peptides with a Net MFI intermediate between the values 92.3 and 63.1, obtained with the two positive control peptides, and 12 peptides with an Net MFI ranging between 29.6 and 63. Fifteen of the in silico predicted peptides (27.2 %) did not confer stabilization to the A2 molecules, showing a Net MFI lower than 14, obtained with the HepB negative control peptide.

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Example 7

Some HLA-A2 binders are recognized by CD8⁺ T cells from DNA-immunized transgenic mice

The *in vitro* assay with RMA-S/A2 cells allowed the definition of a set of peptides which were able to bind and stabilize the HLA-A2 molecules on the cell surface. To gain information about the possibility that the predicted epitopes could indeed be generated by *in vivo* processing of the antigens from which they were derived, peptide recognition by CD8⁺ T cells was studied under conditions in which the related complete antigen was intracellularly expressed and presented *in vivo*. The full-length ORF sequences coding for 13 *Chlamydia*1 proteins, including a total of 24 predicted peptides, were cloned into a suitable DNA expression vector and each recombinant plasmids was used to immunize distinct groups of transgenic mice expressing a chimeric class I molecule composed of the α1 and α2 domains of HLA-0201 and the α3 domains, transmembrane and cytoplasmic, of H-2K^b [35].

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The ORF sequences were selected among those containing either one or more epitopes positive in the *in vitro* assay or a combination of positive and negative epitopes. The ORF sequence corresponding to the outer membrane protein A (OMPA, CPn 0695) was included in this analysis, since human MHC-I-restricted epitopes have already been reported for this protein in *C. trachomatis* [18;36]. One coding sequence, related to gene CPn 0131 was chosen, which included four epitopes, all negative in the *in vitro* stabilization assay. After three immunization cycles, transgenic mice were sacrificed, spleen CD8⁺ T cells were isolated, stimulated for 20 hour with the corresponding peptide and *ex vivo* IFN-γ production was assessed using an enzymelinked immunospot (ELISpot) assay.

Results 7

DNA-mediated expression of the ORFs including peptides CH-6 (CPn 0811), CH-7 (CPn 0623), CH-10 (CPn 0828), CH-13 (CPn 0695, OMPA) and CH-37 (CPn 0210) were associated with numbers of spot forming cells (SFC) significantly higher than those obtained with the HepB unrelated peptide, whereas some peptides related to antigens coded by genes CPn 0131, CPn 0323 and CPn 0062 induced SFC values only 2-3 times higher than the HepB control peptide (Table 5). Peptides related to antigens coded by genes CPn 0132, CPn 0322, CPn 0325, CPn 0415 and CPn 0728 did not induce any IFN-y production (data not shown).

Example 8

To test the capacity of peptides to amplify specific CD8⁺ T cell populations *in vitro*, some of these plasmids were used to repeat the DNA immunization experiment and to determine by flow cytometry the intracellular IFN-γ production by CD8⁺ T cells, both *ex vivo* and after a 6 day stimulation in the presence of the relevant peptides. In the attempt to establish a direct correlation between IFN-γ production by CD8⁺ T cells and HLA-A2 specific restriction, the experiment was carried out with both transgenic and non transgenic syngenic mice. The plasmids used contained genes CPn 0695, CPn 0811 and CPn 0823, including peptides CH-13, CH-6 and CH-7 respectively, which were highly positive in the *in vitro* binding and in the ELISpot assays and gene CPn 0323, including six different peptides, all of them with ELISpot values slightly higher than background

Results 8

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The results of the experiment are summarized in Table 6, while representative dot plots from flow cytometric analysis of splenocytes stimulated with peptide CH-6 are shown in Fig. 4. When fresh spleen cells of DNA-immunized transgenic mice were pulsed with the tested peptides, only CH-6 or CH-7 induced relative fold increase (RFI) values about 5 times higher than those obtained pulsing the same cells with the HepB negative control peptide (Table 6, 4.58 and 5.2 RFI respectively).

When short term T cell lines (TCLs) instead of fresh splenocytes were used, a larger panel of peptides were able to trigger a significantly higher IFN-γ production by CD8⁺ T cells (Table 6). In fact, in addition to peptides CH-6 and CH-7, also peptides CH-13, CH-44, CH-45 and CH-46 were recognized by CD8⁺ T cell populations significantly larger than those induced by pulsing the same cells with the HepB peptide (RFI > 5). Importantly, peptide-induced IFN-γ production, appeared to be largely HLA-A2-dependent, since when the same experiments were carried out with non transgenic mice, the RFI values obtained were reliably lower (Table 6). The fact that non transgenic and transgenic spleen cells were both efficiently activated using the polyclonal stimulus (anti-CD3/anti-CD28), reinforced the hypothesis that the lower CD8⁺ T cells induction in non transgenic mice was due to the absence of specific interactions between the peptides and the human HLA-A2 molecules.

CD8⁺ T cells of transgenic mice infected with *C. pneumoniae* recognize HLA-A2 binders in vivo

It has been recently shown that infection of mice with C. pneumoniae elicits a pathogen-specific murine class I-restricted immune response [22]. Therefore, we asked whether any of the A2 in vitro binders could be recognized by specific CD8⁺ T cells that are clonally selected during the immune response raised against the corresponding native antigen in C. pneumoniae infected cells.

To address this issue, HLA-A2 transgenic mice were intranasally infected with a non 30 lethal dose of C. pneumoniae EBs and challenged with an equal dose of bacteria one month later, before being sacrificed to obtain splenocytes that were used to measure IFN-γ production by CD8⁺ T cells. Since no appreciable IFN-γ-production could be observed if splenocytes from infected mice were tested directly ex vivo (data not shown), spleen cells were cultured with each individual peptide or with the HepB 35 irrelevant peptide for 6 days. The resulting short-term TCLs were then pulsed again for 6 hours with the same peptides and intracellular IFN-γ production by CD8⁺ T cells was assessed. The results obtained with 40 tested peptides are shown in Fig. 5A. Sixteen peptides (CH-2, CH-7, CH-8, CH-10, CH-13, CH-15, CH-20, CH-21, CH-28, CH-35, CH-37, CH-45, CH-46, CH-47, CH-50 and CH-55) elicited the strongest 40 CD8⁺ responses (1 to 7.1 % of IFN-γ-producing CD8⁺ T cells), while 19 peptides elicited low but consistent responses (percentages of CD8+/IFN-γ+ T cells between 0.3 and 0.9). Five peptides did not induce percentages of IFN-γ-producing CD8+ T cells significantly higher than those observed in response to the HepB control peptide. 45

When eight among the most reactive peptides were used again to pulse splenocytes of both transgenic and non transgenic mice infected with C. pneumoniae, seven of them were recognized by specific $CD8^+/IFN-\gamma^+$ T cell populations present only in the transgenic mice, while peptide CH-7 was recognized by $CD8^+$ T cells present in both mice groups (Fig. 5B).

General Discussion of Results in Examples 6-8

In this work we have described peptides derived from *C. pneumoniae* antigens identified as putative T cell epitopes recognized by the common human class I MHC A2 haplotype.

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Understanding C. pneumoniae-specific CD8⁺ T cell-mediated immune response and designing protective vaccines rely on the possibility of identifying bacterial antigens or epitopes recognized by CD8⁺ T cells. Whereas the induction of a CTL-dependent immune response is predictable in response to pathogens which replicate in the cellular cytosol, providing antigens which can enter the cellular MHC-I presentation pathway, in the case of Chlamydiae it is not immediately obvious which antigens are made available to the proteasome and how they reach the cytosol, since these bacteria have a stringent intravacuolar localization inside the infected cell.

We have chosen an *in vivo* system based on HLA-A2 transgenic mice to test which of the predicted peptides could be recognized by specific CD8⁺ T cells following either DNA immunization with individual antigen coding genes or infection with *C. pneumoniae*. Our choice of a murine model is supported by previously published data. Wizel *et al.* [22], recently reported the first evidence that CD8⁺ T cells specific for *C. pneumoniae* antigens are induced in infected mice, and identified bacterial-derived murine MHC-I-restricted T cell epitopes able to trigger either lysis of *C. pneumoniae* infected cells or *in vitro* inhibition of the pathogen intracellular growth. These findings seem to confirm that some *C. pneumoniae* antigens can indeed reach the cytosol of infected cells and enter the MHC-I presentation pathway, i.e. during remodeling that occurs during *Chlamydia* replication or following autolysis of developing bacterial particles [22].

Furthermore, Kuon et al. [42] recently reported the identification of 11 C. trachomatis-derived HLA-B27-restricted peptides, capable of stimulating CD8⁺ T cells obtained from patients with Chlamydia-induced reactive arthritis. Importantly, 8 of them overlapped those selected by analyzing splenocytes of HLA-B27 transgenic mice infected with C. trachomatis, indicating that antigen processing can be closely reproduced using the murine animal model, although differences between murine and human antigen processing and T cell repertoires have been hypothesized [43].

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The experiment which we have performed with *C. pneumoniae* infected A2 transgenic mice revealed that at least 16 peptides were recognized by IFN-γ-positive CD8⁺ T cell populations, which were actually expanded as a consequence of bacterial infection, since we could not detect IFN-γ production pulsing spleen cells from non infected transgenic mice with the same peptides (data not shown). These results suggest that the corresponding *Chlamydial* antigen may be able to enter the MHC-I presentation pathway. The finding that a number of these peptides can also be recognized by specific CD8⁺ T cells when the corresponding protein is separately expressed by DNA immunization, strongly reinforces the hypothesis that the responses observed with the infected mice are indeed specific for the *in silico* predicted peptide epitopes and their corresponding antigens. Importantly, the comparisons of peptide-induced IFN-γ-positive CD8⁺ T cells in A2 transgenic and non transgenic mice, either immunized with DNA or infected with *C. pneumoniae*, indicate that this recognition event is largely A2-specific.

Though, we cannot rule out the possibility that some of the tested peptides are also able to interact with the murine class-I MHC molecules, as suggested by the result obtained with CH-7 in infected non transgenic mice (Fig. 5) and by the RFI values obtained with CH-7, CH-8 and CH-13 in DNA-immunized non transgenic mice (Table 6).

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Both with DNA immunization and bacterial infection, we were able to show that the OMPA-derived CH-13 peptide induces a specific CD8⁺ T cell response in A2 transgenic mice. These results appear to validate the choice of this animal model, 10 since our observation that OMPA can enter the MHC-I presentation pathway correlates with the previous identification of HLA-A2-restricted and of murine MHC-I-restricted epitopes in OMPA proteins of C. trachomatis [18] and of C. pneumoniae [23] respectively. With the exception of CH-13 and CH-17, all the other peptides recognized by CD8⁺ T cells of infected mice belong to C. pneumoniae antigens for 15 which neither human nor murine T cell epitopes have been identified [22;23]. Interestingly, a couple of positively reacting peptides (CH-50 and CH-55) belong to the group of polymorphic outer membrane proteins [44;45], while most of the others are part of the group of Type III secretion system-related proteins [45;46]. Peptides CH-7 and CH-8, both included in protein T of the Yersinia outer protein (Yop) 20 system [47] and CH-10, included in protein J, which is part of the same translocation system, appear to be particularly reactive in the assay with the infected mice (Fig.

This is also true for other peptides included in antigens which are again related to the type III secretion system, such as CH-45, CH-46, and CH-47, all present in the low calcium response protein D. Intriguingly, CH-8, which is the most reactive peptide in the assay with the infected mice, does not seem to be recognized by a specific T cell population when the corresponding antigen is expressed by DNA immunization (Tables 5 and 6). This may be due to different factors, i.e. low *in vivo* expression level of the injected DNA or altered protein conformation.

On the other hand, we should also consider the possibility that, following infection of mice with C. pneumoniae, this peptide is recognized by a $CD8^+$ T cell population which is instead specific for an epitope derived from an unidentified C. pneumoniae antigen having a closely related sequence. Contrarily to CH-8, stimulation of spleen cells from infected transgenic mice with peptide CH-6 did not allow the detection of IFN- γ^+ /CD8 $^+$ T cells (Fig. 5A), but the same peptide was clearly reactive in the DNA immunization experiments (Tables 5 and 6). This may suggest that Low Calcium Response Protein H is not available for the cellular proteasome, but we could also assume either that the amount of peptide available to the MHC-presenting machinery is not sufficient to induce a cell response which is detectable with our assay, or that the reacting CD8 $^+$ T cell population does not expand over the detection limit of our assay.

On the whole, the results presented here allowed the identification of a number of antigens which may be available for proteasome-mediated processing in the course of C. pneumoniae infections, proposing them as possible targets for a HLA-A2-dependent cellular immune response. Further analysis will be required to verify if the specifically induced CD8⁺ T cells are able to recognize and induce lysis of peptide pulsed or C. pneumoniae infected mammalian cells and, possibly, to correlate the

identified T cell epitopes with CD8⁺ T cell populations naturally induced in C. pneumoniae infected patients. Importantly, the results obtained with DNA-mediated expression of distinct antigens, can represent an initial step towards the definition of a significant set of C. pneumoniae HLA-A2-restricted epitopes, which could eventually be combined in DNA minigenes in the attempt to induce a CTL-dependent anti-Chlamydia protective immune response

Example 9

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Immunizations with Combinations of the First Antigen Group

The five antigens of the first antigen group (OmpH-like protein, pmp10, pmp2, Enolase, OmpH-like, CPn0042 and CPn00795 were prepared as described in the Materials and Methods Section above for Examples 1-4. The antigens are expressed and purified. Compositions of antigen combinations are then prepared comprising five antigens per composition (and containing 15 μg of each antigen per composition).

15 CD1 mice are divided into seven groups (5-6 mice per group for groups 1 through 4; 3 to 4 mice for groups 5, 6 and 7), and immunized as follows:

Group	Immunizing Composition	Route of Delivery
1.	Mixture of 5 antigens (15 μg/each) + CFA	Intra-peritoneal
2	Mixture of 5 antigens (15 μg/each) +AlOH (200μg)	Intra-peritoneal
3	Mixture of 5 antigens (15 μg/each) + AlOH (200μg) + CpG (10μg)	Intra-peritoneal
4	Complete Freunds Adjuvant (CFA)	Intra-peritoneal
5	Mixture of 5 antigens (5 μg/each) + LTK63 (5μg)	Intranasal
6	AlOH (200μg) + CpG (10μg)	Intra-peritoneal
7	LTK63 (5µg)	Intranasal

Mice are immunized at two week intervals. Two weeks after the last immunization, all mice are challenged by intravaginal infection with *Chlamydia pneumoniae* serovar D.

Experiment 9 was repeated with another group of CPn antigens. These were: CPn0385 (PepA), CPn0324 (LcrE), CPn0503 (DnaK), CPn0525 (Hypothetical) and CPn0482 (ArtJ). These antigens are combined and administered with and without alum and CpG as described in Experiment 9.

Summary

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Applicants have identified a number of CPn proteins with desirable immunological and/or biological properties. Specifically, at least twelve CPn proteins have been identified which are capable of inducing the production of antibodies, which can neutralise, in a dose-dependent manner, the infectivity of *C. pneumoniae* in *in vitro* cell cultures. The induction of neutralising antibodies is important because it prevents infectious EBs from invading human tissues. Furthermore, at least six of these CPn proteins were also capable of attenuating *Chlamydial* (*C. pneumoniae*) infection in a *in vivo* hamster model. In addition, some of these CPn proteins were also capable of inducing not only adequate T-cell responses but also high serum levels of neutralising antibodies.

Apart from very recent unpublished results on pmp21, this is the first time that antisera to recombinant pmps (pmp2 and pmp10) are reported to have neutralising properties. Interestingly, whilst antiserum against CPn0525 gave negative in vitro results (ie no 15 neutralising activity), the CPn0525 protein gave 97 per cent protection from spleen infection in an in vivo hamster immunisation assay (see Table 2) (ie a positive in vivo result). Likewise, whilst antiserum against Cpn0498 gave negative in vitro results (ie no neutralising activity), the CPn0498 protein gave 94 per cent protection from spleen infection in an in vivo hamster immunisation assay (ie a positive in vivo result). Thus a 20 positive signal obtained in the FACS assay does not guarantee a corresponding positive in vitro neutralization activity and conversely a negative neutralization activity does not mean that a positive in vivo result can be excluded (also look for an example that a negative FACS assay result for a given antibody does not mean that a 25 neutralizing activity can be excluded.

Some of the results obtained by screening the panel of recombinant antigens with the *C.pneumoniae in vitro* neutralization assay are shown in Table 2. Just by a cursory look at the 'current annotation' column it can be seen that both in Table 1 and 2 are listed antigens, like the members of the family of heterogeneous polymorphic membrane proteins (PMP), which, on the basis of published literature data, could be reasonably expected to be surface-exposed and possibly induce neutralizing antibodies, but there are also proteins which could be considered so far only hypothetical, and proteins which just on the basis of their current functional annotation could not be at all expected to be found on the bacterial surface.

The characterisation for the first time of some of these CPn proteins in terms of not only neutralising properties but also different score profiles in a panel of screening tests is an important contribution to the art because it facilitates the selective combination of CPn antigens with particular immunological and biological properties.

In conclusion, this paper describes a group of recombinant antigens which can induce antibodies inhibiting the infectivity of *C. pneumoniae in vitro* and have protective effects *in vivo*..

All publications mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described methods and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the

invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in molecular biology or related fields are intended to be covered by the present invention.

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Figure 1A

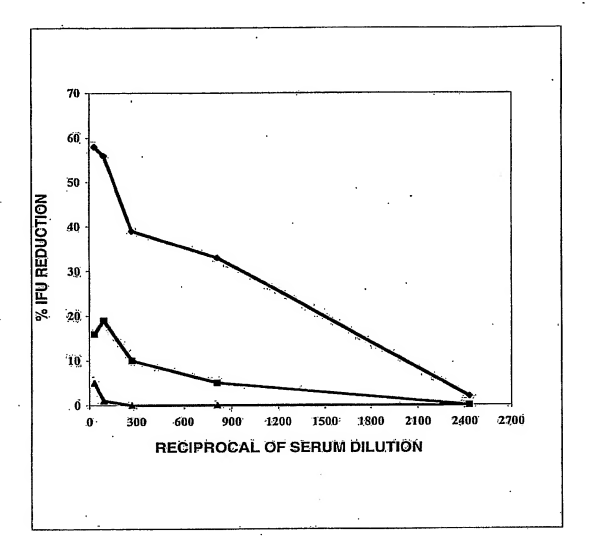
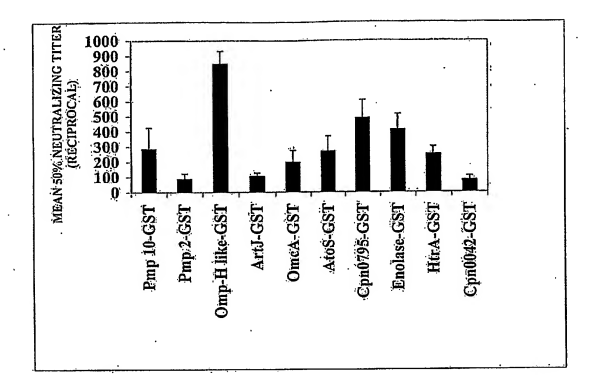


Figure 1B



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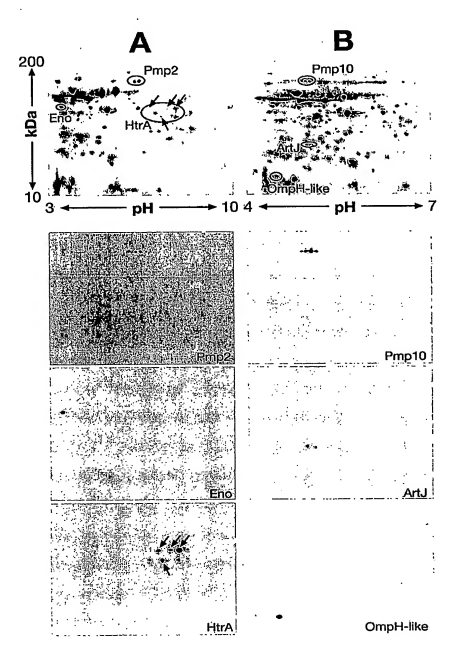


Figure 2

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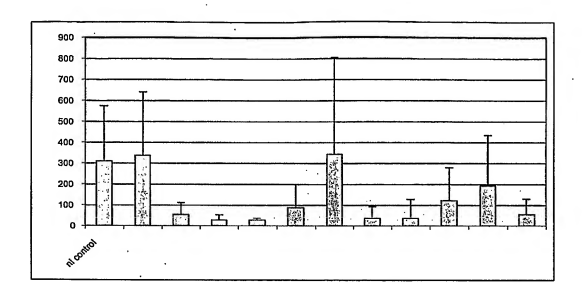


Figure 3. Mean numbers of *C.pneumoniae* IFU recovered from equivalent spleen samples from immunized and mock-immunized hamsters following a systemic challenge. Standard deviation values are shown above the bars. Antigens which induced significant protection are highlighted with an asterisk above the corresponding bar. All antigens were were delivered in Freund's adjuvant. n.i. = non immunized controls

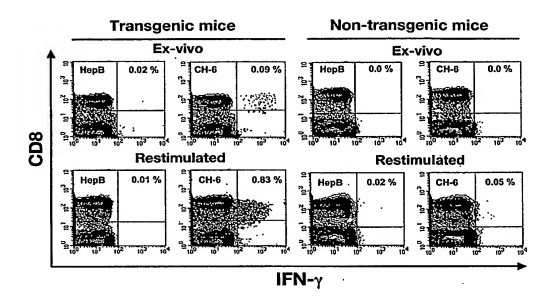
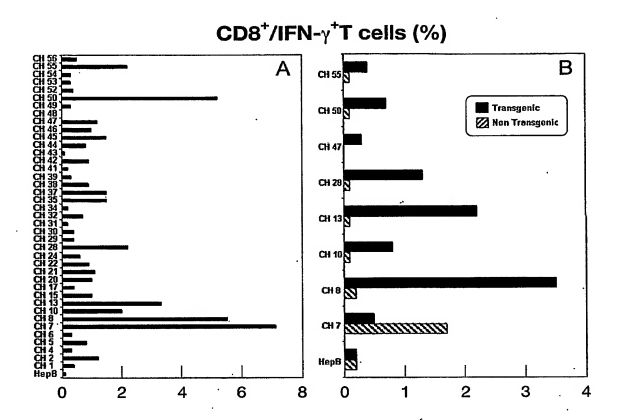


Figure 5



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EXPRESS MAIL NO .: EL 987 060 923 US

APPLICATION DATA SHEET

Application Information

To Be Assigned Application number:

March 2, 2004 Filing Date:

Application Type: **Provisional**

Subject Matter: Utility

To Be Assigned Suggested classification:

Suggested Group Art Unit: To Be Assigned

CD-ROM or CD-R? None

Number of CD disks:

Number of copies of CDs:

Sequence submission? None

Computer Readable Form (CRF)? None

Number of copies of CRF:

Immunogenic Compositions For Chlamydia Title:

No

Pneunomiae

21431.001 Attorney Docket Number:

Request for Early Publication?

No Request for Non-Publication?

Suggested Drawing Figure:

Total Drawing Sheets: 6

No Small Entity?

No 1 Petition included?

Petition Type:

Licensed U.S. Gov't Agency: No

Contract or Grant No:

No Secrecy Order in Parent Appl.?

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First Applicant Information

	•
Applicant Authority Type:	Inventor
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Status:	Full Capacity
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Middle Name: .	
Family Name:	Grandi
Name Suffix:	
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Postal or Zip Code of mailing address:	94662-8097
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Primary Citizenship Country:	
Status:	Full Capacity
Given Name:	Giulio
Middle Name:	
Family Name:	Ratti
Name Suffix:	
City of Residence:	

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State or Province of Residence:

Country of Reside	nce:		•
Street of mailing a	ddress:	c/o Chiron Corporatio	n, P.O. Box 8097
City of mailing add	iress:	Emeryville	
State or Province	of mailing address:	CA	
Country of mailing	address:	US	
Postal or Zip Code	e of mailing address:	94662-8097	
Correspondence	Information		
Correspondence (Customer Number:	27476	•
Street of mailing a	ddress:		
City of mailing add	dress:		
State or Province	of mailing address:		
Country of mailing	address:		
Postal or Zip Code	e of mailing address:		
Phone number:		,	•
Fax Number:		•	
E-Mail address:			
Representative I	nformation		
Representative C	Customer Number:		
Domestic Priority	y Information		
Application :	Continuity Type:	Parent Application:	Parent Filing Date:
		•	

Application :	Continuity Type:	Parent Application:	Parent Filing Date:
			_

Foreign Priority Information

Country:	Application number:	Filing Date:	Priority Claimed:
		·	·
·		·	

Assignee Information

Assignee name:	Chiron S.r.l.
Street of mailing address:	4560 Horton Street
City of mailing address:	Emeryville
State or Province of mailing address:	CA
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